



Figure 1: 254P1D6B SSH sequence of 186 nucleotides (SEQ ID NO: 1).

1 GATCCACAGA TAGGACACAA TTCTTTGGTC ATCAGTAGAC CTTGAACCAT CCAAAGTAAT  
61 GGAATTATTG GGAAGCACAA GAACATGTCT GCCACCAGCC CGGGCTCTGG GAGGACTATT  
121 ATTTTCCTTC TTCACAGCCA CAGTGAGGGT GGACGTGCTG CTCAGTCCCT GCTGGTCTTT  
181 TACTGTCAAA CGGAAGTGGT AGGTCCCCAC CTGGAGACCA GTCACAGTGG CTATTGCTTT  
241 GTCAATATTT TCCATCTCCA CTGCACTGGG GCCTCTGACG TGCT

Figure 2:

Figure 2A. The cDNA (SEQ ID NO. : 2) and amino acid sequence (SEQ ID NO. : 3) of 254P1D6B v.1 clone LCP-3.

The start methionine is underlined. The open reading frame extends from nucleic acid 512-3730 including the stop codon.

```
1 gctgccgcgggcgggtggggcggggatccccgggggtgcaaccttgctccacctgtgctgc
61 cctcggcgggcctggctggccccgcgcagagcggcgggcgctcgctgtcactgccgga
121 ggtgagagcgcagcagtagcttcagcctgtcttgggcttggtccagattcgctcctctgg
181 ggctacgtcccggggaagaggaagcgaggattttgctgggggtggggctgtacctctaac
241 agcaggtgcgcgcgcgaggggtgtgaactgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtg
301 taagacctgcgatgacgacgaggaggaacaagtgggacggcgagtgtatgctcagggccag
361 cagcaacgcatggggcgagcttcagtgtcgccagcagtgaccacagttcttgaggccaaa
421 tctggctcctaaaaaacatcaaaggaagcttgaccaaactctcttcagggccgcctcag
1 M A P P T G V L S S
481 aagcctgccatcaccactgtgtggtgcacaATGGCGCCCCCACAGGTGTGCTCTCTTC
11 L L L L V T I A G C A R K Q C S E G R T
541 ATTGCTGCTGCTGGTGACAATTGCAGGTTGTGCCCCGTAAGCAGTGCAGCGAGGGGAGGAC
31 Y S N A V I S P N L E T T R I M R V S H
601 ATATTCCAATGCAGTCATTTACCTAACTTGGAAACCACCAGAATCATGCGGGTGTCTCA
51 T F P V V D C T A A C C D L S S C D L A
661 CACCTTCCCTGTCTGACTGCACGGCCGCTTGTGTGACCTGTCCAGCTGTGACCTGGC
71 W W F E G R C Y L V S C P H K E N C E P
721 CTGGTGGTTTCGAGGGCCGCTGTACCTGGTGAGCTGCCCCACAAAGAGAACTGTGAGCC
91 K K M G P I R S Y L T F V L R P V Q R P
781 CAAGAAGATGGGCCCCATCAGGTCTTATCTCACTTTTGTGCTCCGGCCTGTTCAGAGGCC
111 A Q L L D Y G D M M L N R G S P S G I W
841 TGCACAGCTGCTGGACTATGGGGACATGATGCTGAACAGGGGCTCCCCCTCGGGGATCTG
131 G D S P E D I R K D L P F L G K D W G L
901 GGGGACTCACCTGAGGATATCAGAAAGGACTTGMCCCTTCTAGGCAAAGATTGGGGCCT
151 E E M S E Y S D D Y R E L E K D L L Q P
961 AGAGGAGATGTCTGAGTACKCAGATGACTACCGGGAGCTGGAGAAGGACCTCTTGCAACC
171 S G K Q E P R G S A E Y T D W G L L P G
1021 CAGTGGCAAGCAGGAGCCCAGAGGGAGTGCCGAGTACACGGACTGGGGCCTACTGCCGGG
191 S E G A F N S S V G D S P A V P A E T Q
1081 CAGCGAGGGGGCCTTCAACTCCTCTGTTGGAGACAGTCTGCGGTGCCAGCGGAGACGCA
211 Q D P E L H Y L N E S A S T P A P K L P
1141 GCAGGACCCTGAGCTCCATTACCTGAATGAGTCGGCTTCAACCCCTGCCCCAAACTCCC
231 E R S V L L P L P T T P S S G E V L E K
1201 TGAGAGAAGTGTGTTGCTTCCCTTGCCGACTACTCCATCTTCAGGAGAGGTGTTGGAGAA
251 E K A S Q L Q E Q S S N S S G K E V L M
1261 AGAAAAGGCTTCTCAGCTCCAGGAACAATCCAGCAACAGCTCTGGAAGAGAGGTTCTAAT
271 P S H S L P P A S L E L S S V T V E K S
1321 GCCTTCCCATAGTCTTCTCCGGCAAGCCTGGAGCTCAGCTCAGTCACCGTGGAGAAAAG
```

291 P V L T V T P G S T E H S I P T P P T S  
1381 CCCAGTGCTCACAGTCACCCCGGGGAGTACAGAGCACAGCATCCCAACACCTCCCCTAG  
311 A A P S E S T P S E L P I S P T T A P R  
1441 CGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATATCTCCTACCACTGCTCCCAG  
331 T V K E L T V S A G D N L I I T L P D N  
1501 GACAGTGAAAGAACTTACGGTATCGGCTGGAGATAACCTAATTATAACTTTACCCGACAA  
351 E V E L K A F V A P A P P V E T T Y N Y  
1561 TGAAGTTGAACTGAAGGCCTTTGTTGCGCCAGCGCCACCTGTAGAAACAACCTACAATA  
371 E W N L I S H P T D Y Q G E I K Q G H K  
1621 TGAATGGAATTTAATAAGCCACCCACAGACTACCAAGGTGAAATAAAACAAGGACACAA  
391 Q T L N L S Q L S V G L Y V F K V T V S  
1681 GCAAACCTCTTAACCTCTCTCAATTGTCCGTCGGACTTTATGTCTTCAAAGTCACTGTTTC  
411 S E N A F G E G F V N V T V K P A R R V  
1741 TAGTGAAAACGCCTTTGGAGAAGGATTTGTCAATGTCACTGTTAAGCCTGCCAGAAGAGT  
431 N L P P V A V V S P Q L Q E L T L P L T  
1801 CAACCTGCCACCTGTAGCAGTTGTTTCTCCCCAACTGCAAGAGCTCACTTTGCCTTTGAC  
451 S A L I D G S Q S T D D T E I V S Y H W  
1861 GTCAGCCCTCATTGATGGCAGCCAAAGTACAGATGATACTGAAATAGTGAGTTATCATTG  
471 E E I N G P F I E E K T S V D S P V L R  
1921 GGAAGAAATAAACGGGCCCTTCATAGAAGAGAAGACTTCAGTTGACTCTCCCGTCTTACG  
491 L S N L D P G N Y S F R L T V T D S D G  
1981 CTTGTCTAACCTTGATCCTGGTAACTATAGTTTCAGGTTGACTGTTACAGACTCGGACGG  
511 A T N S T T A A L I V N N A V D Y P P V  
2041 AGCCACTAACTCTACAACCTGCAGCCCTAATAGTGAACAATGCTGTGGACTACCCACCAGT  
531 A N A G P N H T I T L P Q N S I T L N G  
2101 TGCTAATGCAGGACCAAATCACACCATAACTTTGCCCCAAACTCCATCACTTTGAATGG  
551 N Q S S D D H Q I V L Y E W S L G P G S  
2161 AAACCAGAGCAGTGACGATCACCAGATTGTCTCTATGAGTGGTCCCTGGGTCTGGGAG  
571 E G K H V V M Q G V Q T P Y L H L S A M  
2221 TGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCATACCTTCATTTATCTGCAAT  
591 Q E G D Y T F Q L K V T D S S R Q Q S T  
2281 GCAGGAAGGAGATTATACATTTTCACTGAAGGTGACAGATTCTTCAAGGCAACAGTCTAC  
611 A V V T V I V Q P E N N R P P V A V A G  
2341 TGCTGTRGTGACTGTGATTGTCCAGCCTGAAAACAATAGACCTCCAGTGGCTGTGGCCGG  
631 P D K E L I F P V E S A T L D G S S S S  
2401 CCCTGATAAAGAGCTGATCTTCCCAGTGGAAAGTGCTACCCTGGATGGGAGCAGCAGCAG  
651 D D H G I V F Y H W E H V R G P S A V E  
2461 CGATGACCACGGCATTGTCTTCTACCACTGGGAGCACGTCAGAGGCCCACTGCAGTGGAA  
671 M E N I D K A I A T V T G L Q V G T Y H  
2521 GATGGAAAATATTGACAAAGCAATAGCCACTGTGACTGGTCTCCAGGTGGGGACCTACCA  
691 F R L T V K D Q Q G L S S T S T L T V A  
2581 CTTCCGTTTGACAGTGAAAGACCAGCAGGGACTGAGCAGCACGTCCACCCTCACTGTGGC

711 V K K E N N S P P R A R A G G R H V L V  
2641 TGTGAAGAAGGAAAATAATAGTCCTCCAGAGCCCGGGCTGGTGGCAGACATGTTCTTGT  
731 L P N N S I T L D G S R S T D D Q R I V  
2701 GCTTCCCAATAATTCCATTACTTTGGATGGTTCAAGGTCTACTGATGACCAAAGAATTGT  
751 S Y L W I R D G Q S P A A G D V I D G S  
2761 GTCCTATCTGTGGATCCGGGATGGCCAGAGTCCAGCAGCTGGAGATGTCATCGATGGCTC  
771 D H S V A L Q L T N L V E G V Y T F H L  
2821 TGACCACAGTGTGGCTCTGCAGCTTACGAATCTGGTGGAGGGGTGTACACTTTCCACTT  
791 R V T D S Q G A S D T D T A T V E V Q P  
2881 GCGAGTCACCGACAGTCAGGGGGCCTCGGACACAGACACTGCCACTGTGGAAGTGCAGCC  
811 D P R K S G L V E L T L Q V G V G Q L T  
2941 AGACCCTAGGAAGAGTGGCCTGGTGGAGCTGACCCTGCAGGTTGGTGTGGGCAGCTGAC  
831 E Q R K D T L V R Q L A V L L N V L D S  
3001 AGAGCAGCGGAAGGACACCCTTGTGAGGCAGCTGGCTGTGCTGCTGAACGTGCTGGACTC  
851 D I K V Q K I R A H S D L S T V I V F Y  
3061 GGACATTAAGGTCCAGAAGATTCGGGCCCCACTCGGATCTCAGCACCGTGATTGTGTTT  
871 V Q S R P P F K V L K A A E V A R N L H  
3121 TGTACAGAGCAGGCCGCCTTTCAAGGTTCTCAAAGCTGCTGAAGTGGCCCCGAAATCTGCA  
891 M R L S K E K A D F L L F K V L R V D T  
3181 CATGCGGCTCTCAAAGGAGAAGGCTGACTTCTTGCTTTTCAAGGTCTTGAGGGTTGATAC  
911 A G C L L K C S G H G H C D P L T K R C  
3241 AGCAGGTTGCCTTCTGAAGTGTCTGGCCATGGTCACTGCGACCCCTCACAAAGCGCTG  
931 I C S H L W M E N L I Q R Y I W D G E S  
3301 CATTGCTCTCACTTATGGATGGAGAACCTTATACAGCGTTATATCTGGGATGGAGAGAG  
951 N C E W S I F Y V T V L A F T L I V L T  
3361 CAACTGTGAGTGGAGTATATTCTATGTGACAGTGTGGCTTTTACTCTTATTGTGCTAAC  
971 G G F T W L C I C C C K R Q K R T K I R  
3421 AGGAGGTTTCACTTGGCTTTGCATCTGCTGCTGCAAAAGACAAAAAGGACTAAAATCAG  
991 K K T K Y T I L D N M D E Q E R M E L R  
3481 GAAAAAACAAAGTACACCATCCTGGATAACATGGATGAACAGGAAAGAATGGAAGTGAAG  
1011 P K Y G I K H R S T E H N S S L M V S E  
3541 GCCCAAATATGGTATCAAGCACCGAAGCACAGAGCACAACTCCAGCCTGATGGTATCCGA  
1031 S E F D S D Q D T I F S R E K M E R G N  
3601 GTCTGAGTTTGACAGTGACCAGGACACAATCTTCAGCCGAGAAAAGATGGAGAGAGGGAA  
1051 P K V S M N G S I R N G A S F S Y C S K  
3661 TCCAAAGGTTTCCATGAATGGTTCCATCAGAAATGGAGCTTCCTTCAGTTATTGCTCAAA  
1071 D R \*  
3721 GGACAGATAATggcgcagttcattgtaaagtgaaggacccyttgaatccargaccagtc  
3781 agtgggagttacagcacaaaaccactcttttagaatagttcattgaccttcttccccag  
3841 tgggttagatgtgtatccccacgtactaaaagaccggtttttgaaggcacaaaacaaaa  
3901 ctttctcttttaactgagatgcttgtaataagaaataaaggctgggtaaaactytaaag  
3961 tatatacttaaaagagttttgagttttgtagctggcacaaatctcatattaaagatgaac

4021 aacgatttctatctgtagaaccttagagaaggatgaatgaaacaagggttttaaaaagggat  
4081 gatttctgtcttagcygctgtgattgcctctaaggaacagcattctaaacacggtttctc  
4141 ttgtaggacctgcagtcagatggctgtgtatgttaaaatagcttgtctaaggagcacggg  
4201 ccatctgtggaggtagcgagtccttgcatgtagcaagctttctgtgtgacggcaacactc  
4261 gcacagtgccaaagccctcctgggttttaattctgtgctatgtcaatggcagttttcatct  
4321 ctctcaagaaagcagctgttgccattcaagagctaaggaagaatcgatttctaaggact  
4381 gaggcaatagaaagggaggaggagcttaatgcctgcaggttgaaggtagcattgtaac  
4441 attatcttttcttctctaagaaaaactacactgactcctctcgggtgtgttagcagta  
4501 tagttctctaattgtaaacrgatccccagtttacattaartgcaatagaagtgattaattc  
4561 attaaacatttattatgttctgtaggctgtgcgtttggactgccatagataggkataacg  
4621 actcagcaattgtgtatatattccaaaactctgaaatacagtcagtccttaacttggatgg  
4681 cgtgggttatgatactctgggtccccgacaggtaactttccaaaataacttgacatagatgta  
4741 ttcacttcatatgtttaaaaatacatttaagttttctaccgaataaatcttattttcaaa  
4801 catgaaagacaattaaaacattcccaccacaaaagcagtaactcccgagcaattaactgga  
4861 gttaattgtagcctgctacgttgactgggtcagggtagttccccatccacccttggctcct  
4921 gaggtggtggccttgggtgggtgcccttggcattttttgtgggaagattagaatgagagat  
4981 agaaccagtgttgtggtaccaagtgtgagcacacctaacaatctcgttgcacaatgc  
5041 ttttttaacacatgggaaaactaggaatgcattgctgatgaagaagcaaggattttaaac  
5101 accagggcaggagtgccagagaaaatgtttcccatgggttcttaaaaaaattcagctt  
5161 ttaggtgcttttgtcatctccgsagattcatcctcatgggaccatcttatttttactt  
5221 attgtaatttactggggaaaggcagaactaaaaagtgtgtcattttatttttaaaataat  
5281 tgctttgcttatgcctacactttctgtataactagccaattcaatactgtctatagtgtt  
5341 agaaggaaaatgtgattttttttttaaccagtattgagcttcataagcctagaatctg  
5401 ccttatcagggtgaccaggttatggttgtttgcatgcaaagtgaatttctggcataggg  
5461 gacagcagcccaaagttaaagtcacggtcgtaagtgaagaaggagtgaaacatttac  
5521 cgctttakgtacataacatatgcagtttacatactcatttgatccttataatcaaccttg  
5581 aagaggagatactatcattcttatgttgcagatagccctctgaaggcccagagaggttaa  
5641 rtaacttcccagaggtcatggccaagaagttagtggtccaagaactgaatgcaaatTTTT  
5701 taaactgtagagttctgctttccactaaacaaagaactcctgccttgatggatggagggc  
5761 aaattctggtggaacttttgggccacctgaaagtctattcccaggactaagaggaattt  
5821 cttttaatggatccagagagccaaggtcagaggagagatggcctgcatagtctcctgtg  
5881 gatcacacccgggccacccctccctctaggtttacagtggacttcttctgccccctcctcc  
5941 ttttctgtccttggccatctcagcctggcctctctgatccttccatcacagaaggatctt  
6001 gaatctctgggaaatcaaacatcacagtagtgatcagaaagtgagtcctgtcttgtcacc  
6061 ccatttctcatcagaacaaagcacgagatggaatgaccaaccagcattcttcatggtgga  
6121 ctgcttatcattgaggatctttgggagataaagcacgctaagagctctggacagagaaaa  
6181 acaggccctagaatatgggagtggtgtttgtagggtcayargtaacaagcacttttag  
6241 ttgctgggttacattcaatgaaggaggattcatacccatggcattacaaggctaagcatg  
6301 tgtatgactaaggaaactatctgaaaaacatgcagcaaggtaagaaaatgtaccactcaac  
6361 aagccagtgtatgccaccttttgtgcgcggggaggagagtgactaccattgtttttgtgt  
6421 gacaaagctatcatggactatttttaattcttgggttttattgcttaaaatatattttttc  
6481 cctatgtgttgacaaggatatttctaataatcacactattaaatatatgcactaatctaaat

6541 aaaggtgtctgtatcttctgtaatgcttatttttaggggaaattgttttctttatgct  
6601 tcagggtagagggattcccttgagtataggtcagcaaactctggcctgcagcctgtgtgt  
6661 gcacgccccatgagccgaaaagtgggtcttatgttttcaaagtgttaaaaataaataaaa  
6721 aaatttgaaacatgtgaactatatgacattcagatttgtgttcataaataaagttttatt  
6781 ggaacatatcc

**Figure 2B. The cDNA (SEQ ID NO. : 4) and amino acid sequence (SEQ ID NO. : 5) of 254P1D6B v.2. The start methionine is underlined. The open reading frame extends from nucleic acid 512-3730 including the stop codon.**

1 gctgccgcgggcgggtgggcggggatccccgggggtgcaaccttgctccacctgtgctgc  
61 cctcggcgggcctggctggccccgcgcagagcggcgggcgctcgctgtcactgccgga  
121 ggtgagagcgcagcagtagcttcagcctgtcttgggcttgggtccagattcgctcctctgg  
181 ggctacgtccccggggaagaggaagcgaggattttgctgggggtggggctgtacctcttaac  
241 agcaggtgcgcgcgcgaggggtgtgaacgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtg  
301 taagacctgcgatgacgacgaggaggaacaagtgggacggcgagtgatgctcagggccag  
361 cagcaacgcgatggggcgagcttcagtgctgccagcagtgaccacagttcttgaggccaaa  
421 tctggctcctaaaaaacatcaaaggaagcttgaccaaactctcttcaggggccgcctcag  
1 M A P P T G V L S S  
481 aagcctgccatcacccactgtgtggtgcacaATGGCGCCCCCACAGGTGTGCTCTCTTC  
11 L L L L V T I A G C A R K Q C S E G R T  
541 ATTGCTGCTGCTGGTGACAATTGCAGGTTGTGCCCCGTAAGCAGTGCAGCGAGGGGAGGAC  
31 Y S N A V I S P N L E T T R I M R V S H  
601 ATATTCCAATGCAGTCATTTACCTAACTTGGAACCACCAGAATCATGCGGGTGTCTCA  
51 T F P V V D C T A A C C D L S S C D L A  
661 CACCTTCCCTGTCGTAGACTGCACGGCCGCTTGCTGTGACCTGTCCAGCTGTGACCTGGC  
71 W W F E G R C Y L V S C P H K E N C E P  
721 CTGGTGGTTCGAGGGCCGCTGCTACCTGGTGAGCTGCCCCACAAAGAGAACTGTGAGCC  
91 K K M G P I R S Y L T F V L R P V Q R P  
781 CAAGAAGATGGGCCCCATCAGGTCTTATCTCACTTTTGTGCTCCGGCCTGTTCAGAGGCC  
111 A Q L L D Y G D M M L N R G S P S G I W  
841 TGCACAGCTGCTGGACTATGGGGACATGATGCTGAACAGGGGCTCCCCCTCGGGGATCTG  
131 G D S P E D I R K D L P F L G K D W G L  
901 GGGGACTCACCTGAGGATATCAGAAAGGACTTGCCCTTTCTAGGCAAAGATTGGGGCCT  
151 E E M S E Y A D D Y R E L E K D L L Q P  
961 AGAGGAGATGTCTGAGTACGCAGATGACTACCGGGAGCTGGAGAAGGACCTCTTGCAACC  
171 S G K Q E P R G S A E Y T D W G L L P G  
1021 CAGTGGCAAGCAGGAGCCCAGAGGGAGTGCCGAGTACACGGACTGGGGCCTACTGCCGGG  
191 S E G A F N S S V G D S P A V P A E T Q  
1081 CAGCGAGGGGGCCTTCAACTCCTCTGTTGGAGACAGTCCTGCGGTGCCAGCGGAGACGCA  
211 Q D P E L H Y L N E S A S T P A P K L P  
1141 GCAGGACCCTGAGCTCCATTACCTGAATGAGTCGGCTTCAACCCCTGCCCAAACCTCCC  
231 E R S V L L P L P T T P S S G E V L E K

1201 TGAGAGAAGTGTGTTGCTTCCCTTGCCGACTACTCCATCTTCAGGAGAGGTGTTGGAGAA  
251 E K A S Q L Q E Q S S N S S G K E V L M  
1261 AGAAAAGGCTTCTCAGCTCCAGGAACAATCCAGCAACAGCTCTGGAAAAGAGGTTCTAAT  
271 P S H S L P P A S L E L S S V T V E K S  
1321 GCCTTCCCATAGTCTTCCTCCGGCAAGCCTGGAGCTCAGCTCAGTCACCGTGGAGAAAAG  
291 P V L T V T P G S T E H S I P T P P T S  
1381 CCCAGTGCTCACAGTCACCCCGGGGAGTACAGAGCACAGCATCCCAACACCTCCCCTAG  
311 A A P S E S T P S E L P I S P T T A P R  
1441 CGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATATCTCCTACCACTGCTCCCAG  
331 T V K E L T V S A G D N L I I T L P D N  
1501 GACAGTGAAAGAACTTACGGTATCGGCTGGAGATAACCTAATTATAACTTTACCCGACAA  
351 E V E L K A F V A P A P P V E T T Y N Y  
1561 TGAAGTTGAACTGAAGGCCTTTGTTGCGCCAGCGCCACCTGTAGAAACAACCTACAAC  
371 E W N L I S H P T D Y Q G E I K Q G H K  
1621 TGAATGGAATTTAATAAGCCACCCACAGACTACCAAGGTGAAATAAAACAAGGACACAA  
391 Q T L N L S Q L S V G L Y V F K V T V S  
1681 GCAAACCTCTTAACCTCTCTCAATTGTCCGTCGGACTTTATGTCTTCAAAGTCACTGTTTC  
411 S E N A F G E G F V N V T V K P A R R V  
1741 TAGTGAAAACGCCTTTGGAGAAGGATTTGTCAATGTCACTGTTAAGCCTGCCAGAAGAGT  
431 N L P P V A V V S P Q L Q E L T L P L T  
1801 CAACCTGCCACCTGTAGCAGTTGTTTCTCCCCAACTGCAAGAGCTCACTTTGCCTTTGAC  
451 S A L I D G S Q S T D D T E I V S Y H W  
1861 GTCAGCCCTCATTGATGGCAGCCAAAGTACAGATGATACTGAAATAGTGAGTTATCATTG  
471 E E I N G P F I E E K T S V D S P V L R  
1921 GGAAGAAATAAACGGGCCCTTCATAGAAGAGAAGACTTCAGTTGACTCTCCCGTCTTACG  
491 L S N L D P G N Y S F R L T V T D S D G  
1981 CTTGTCTAACCTTGATCCTGGTAACCTATAGTTTCAGGTTGACTGTTACAGACTCGGACGG  
511 A T N S T T A A L I V N N A V D Y P P V  
2041 AGCCACTAACTCTACAACCTGCAGCCCTAATAGTGAACAATGCTGTGGACTACCCACCAGT  
531 A N A G P N H T I T L P Q N S I T L N G  
2101 TGCTAATGCAGGACCAAATCACACCATAACTTTGCCCCAAAACCTCCATCACTTTGAATGG  
551 N Q S S D D H Q I V L Y E W S L G P G S  
2161 AAACCAGAGCAGTGACGATCACCAGATTGTCCTCTATGAGTGGTCCCTGGGTCTGGGAG  
571 E G K H V V M Q G V Q T P Y L H L S A M  
2221 TGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCATACCTTCATTTATCTGCAAT  
591 Q E G D Y T F Q L K V T D S S R Q Q S T  
2281 GCAGGAAGGAGATTATACATTTTCAGCTGAAGGTGACAGATTCTTCAAGGCAACAGTCTAC  
611 A V V T V I V Q P E N N R P P V A V A G  
2341 TGCTGTAGTGAAGTGTGATTGTCCAGCCTGAAAACAATAGACCTCCAGTGGCTGTGGCCGG  
631 P D K E L I F P V E S A T L D G S S S S  
2401 CCCTGATAAAGAGCTGATCTTCCCACTGGAAAGTGCTACCCTGGATGGGAGCAGCAGCAG  
651 D D H G I V F Y H W E H V R G P S A V E

2461 CGATGACCACGGCATTGTCTTCTACCACTGGGAGCACGTCAGAGGCCCCAGTGCAGTGG  
671 M E N I D K A I A T V T G L Q V G T Y H  
2521 GATGGAAAATATTGACAAAGCAATAGCCACTGTGACTGGTCTCCAGGTGGGGACCTACCA  
691 F R L T V K D Q Q G L S S T S T L T V A  
2581 CTTCCGTTTGACAGTGAAAGACCAGCAGGGACTGAGCAGCACGTCCACCCCTACTGTGGC  
711 V K K E N N S P P R A R A G G R H V L V  
2641 TGTGAAGAAGGAAAATAATAGTCCTCCAGAGCCCGGGCTGGTGGCAGACATGTTCTTGT  
731 L P N N S I T L D G S R S T D D Q R I V  
2701 GCTTCCCAATAATTCCATTACTTTGGATGGTTCAAGGTCTACTGATGACCAAAGAATTGT  
751 S Y L W I R D G Q S P A A G D V I D G S  
2761 GTCCTATCTGTGGATCCGGGATGGCCAGAGTCCAGCAGCTGGAGATGTCATCGATGGCTC  
771 D H S V A L Q L T N L V E G V Y T F H L  
2821 TGACCACAGTGTGGCTCTGCAGCTTACGAATCTGGTGGAGGGGGTGTACACTTTCCACTT  
791 R V T D S Q G A S D T D T A T V E V Q P  
2881 GCGAGTCACCGACAGTCAGGGGGCCTCGGACACAGACACTGCCACTGTGGAAGTGCAGCC  
811 D P R K S G L V E L T L Q V G V G Q L T  
2941 AGACCCTAGGAAGAGTGGCCTGGTGGAGCTGACCCTGCAGGTTGGTGTGGGCAGCTGAC  
831 E Q R K D T L V R Q L A V L L N V L D S  
3001 AGAGCAGCGGAAGGACACCCTTGTGAGGCAGCTGGCTGTGCTGCTGAACGTGCTGGACTC  
851 D I K V Q K I R A H S D L S T V I V F Y  
3061 GGACATTAAGGTCCAGAAGATTGGGGCCCACTCGGATCTCAGCACCGTGATTGTGTTTTA  
871 V Q S R P P F K V L K A A E V A R N L H  
3121 TGTACAGAGCAGGCCGCCTTTCAAGGTTCTCAAAGCTGCTGAAGTGGCCCCGAAATCTGCA  
891 M R L S K E K A D F L L F K V L R V D T  
3181 CATGCGGCTCTCAAAGGAGAAGGCTGACTTCTTGCTTTTCAAGGTCTTGAGGGTTGATAC  
911 A G C L L K C S G H G H C D P L T K R C  
3241 AGCAGGTTGCCTTCTGAAGTGTCTGGCCATGGTCACTGCGACCCCTCACAAAGCGCTG  
931 I C S H L W M E N L I Q R Y I W D G E S  
3301 CATTTGCTCTCACTTATGGATGGAGAACCTTATACAGCGTTATATCTGGGATGGAGAGAG  
951 N C E W S I F Y V T V L A F T L I V L T  
3361 CAACTGTGAGTGGAGTATATTCTATGTGACAGTGTGGCTTTTACTCTTATTGTGCTAAC  
971 G G F T W L C I C C C K R Q K R T K I R  
3421 AGGAGGTTTCACTTGGCTTTGCATCTGCTGCTGCAAAAGACAAAAAGGACTAAAATCAG  
991 K K T K Y T I L D N M D E Q E R M E L R  
3481 GAAAAAACAAGTACACCATCCTGGATAACATGGATGAACAGGAAAGAATGGAAGTGAAG  
1011 P K Y G I K H R S T E H N S S L M V S E  
3541 GCCCAAATATGGTATCAAGCACCGAAGCACAGAGCACAACTCCAGCCTGATGGTATCCGA  
1031 S E F D S D Q D T I F S R E K M E R G N  
3601 GTCTGAGTTTGACAGTGACCAGGACACAATCTTCAGCCGAGAAAAGATGGAGAGAGGGAA  
1051 P K V S M N G S I R N G A S F S Y C S K  
3661 TCCAAAGGTTTCCATGAATGGTTCCATCAGAAATGGAGCTTCCTTCAGTTATTGCTCAA  
1071 D R \*



3721 GGACAGATAATggcgcagttcattgtaaagtgaaggaccccttgaatccaagaccagtc  
3781 agtgggagttacagcacaaaaccactcttttagaatagttcattgaccttcttccccag  
3841 tgggttagatgtgtatccccacgtactaaaagaccggtttttgaaggcacaaaacaaaa  
3901 ctttgctcttttaactgagatgcttgtaataagaaataaaggctgggtaaaactctaagg  
3961 tatatacttaaaagagttttgagttttgtagctggcacaatctcatattaaagatgaac  
4021 aacgattttctatctgtagaaccttagagaagggtgaatgaaacaagggttttaaaaaggat  
4081 gatttctgtcttagccgctgtgattgcctctaaggaacagcattctaaacacggtttctc  
4141 ttgtaggacctgcagtcagatggctgtgtatgttaaaatagcttgtctaagaggcacggg  
4201 ccatctgtggagggtacggagctcttgcatgtagcaagctttctgtgctgacggcaacactc  
4261 gcacagtccaagccctcctgggttttaattctgtgctatgtcaatggcagttttcatct  
4321 ctctcaagaaagcagctgttggccattcaagagctaaggaagaatcgtattctaaggact  
4381 gaggcaatagaaaggggaggaggagcttaatgccgtgcagggtgaaggtagcattgtaac  
4441 attatcttttcttctctaagaaaaactacactgactcctctcgggtgttgttttagcagta  
4501 tagttctctaattgtaaaccggatccccagtttacattaaatgcaatagaagtgattaattc  
4561 attaagcatttattatgttctgtaggctgtgcgtttggactgccatagatagggataacg  
4621 actcagcaattgtgtatatattccaaaactctgaaatacagtcagtccttaacttggatgg  
4681 cgtggttatgatactctggtccccgacaggtactttccaaaataacttgacatagatgta  
4741 ttcacttcatatgtttaaaaatacatttaagtttttctaccgaataaatcttatttcaaa  
4801 catgaaagacaattaaaacattcccacccacaaagcagtactcccgagcaattaactgga  
4861 gttaattgtagcctgctacgttgactgggttcagggttagttcccatccacccttggctct  
4921 gaggtcgggtggccttgggtgggtgcccttggcattttttgtgggaagattagaatgagagat  
4981 agaaccagtggtgtggtaccaagtgtgagcacacctaacaatatcctgttgcaaatgc  
5041 ttttttaacacatgggaaaactaggaatgcattgctgatgaagaagcaaggattttaaac  
5101 accagggcaggagtgccagagaaaaatgtttcccatgggttcttaaaaaaaattcagctt  
5161 ttaggtgcttttgtcatctcccgaggtattcatcctcatgggaccatcttatttttactt  
5221 attgtaatttactggggaaaggcagaactaaaaagtgtgtcattttatttttaaaataat  
5281 tgctttgcttatgcctacactttctgtataactagccaattcaatactgtctatagtgtt  
5341 agaaggaaaatgtgatttttttttttaaccagattgagcttcataagcctagaatctg  
5401 ccttatcaggtgaccagggttatgggtgtttgcatgcaaagtgaatttctggcataggg  
5461 gacagcagcccaaatgtaaagtcacgggcgtaatgaggaagaagggtgaacatttac  
5521 cgctttatgtacataacatatgcagtttacatactcatttgatccttataatcaaccttg  
5581 aagaggagatactatcattcttatgttgcatagccctctgaaggcccagagagggttaa  
5641 gtaacttcccagaggtcatggccaagaagttagtggtccaagaactgaatgcaattttt  
5701 taaactgtagagttctgctttccactaaacaaagaactcctgccttgatggatggagggc  
5761 aaattctggtggaacttttgggccacctgaaagttctattcccaggactaagaggaattt  
5821 cttttaatggatccagagagccaagggtcagagggagagatggcctgcatagtctcctgtg  
5881 gatcacacccgggccacccctcctctaggtttacagtggacttcttctgcccctcctcc  
5941 ttttctgtccttggccatctcagcctggcctctctgatccttccatcacagaaggatctt  
6001 gaatctctgggaaatcaaacatcacagtagtgatcagaaaagtgaagtctgtctgtcacc  
6061 ccatttctcatcagaacaaagcacgagatggaatgaccaaccagcattcttcatgggtga  
6121 ctgcttatcattgaggatctttgggagataaagcacgctaagagctctggacagagaaaa  
6181 acaggccctagaatatgggagtggtgtttgtagggctcataggctaacaagcactttag

6241 ttgctggtttacattcaatgaaggaggattcatacccatggcattacaaggctaagcatg  
6301 tgtatgactaaggaactatctgaaaaacatgcagcaaggtaagaaaatgtaccactcaac  
6361 aagccagtgatgccaccttttgtgcgcggggaggagagtgactaccattgttttttgtgt  
6421 gacaaagctatcatggactattttaatcttggttttatttgcttaaaatatattttttc  
6481 cctatgtgttgacaaggatatttctaataatcacactattaaatatatgcactaatctaaat  
6541 aaagggtgtctgtattttctgtaatgcttatttttaggggaaatttgttttctttatgct  
6601 tcagggttagagggttcccttgagtataggtcagcaaacctctggcctgcagcctgtgtgt  
6661 gcacgccccatgagccgaaaagtgggtcttatgttttcaaattgggttaaaaaataaaaa  
6721 aaatttgaaacatgtgaactatatgacattcagatttgtgttcataaataaagttttatt  
6781 ggaacatatcc

**Figure 2C. The cDNA (SEQ ID NO. : 6) and amino acid sequence (SEQ ID NO. : 7) of 254P1D6B v.3. The start methionine is underlined. The open reading frame extends from nucleic acid 739-3930 including the stop codon.**

1 gctgccgcgggcggtgggcggggatccccgggggtgcaaccttgctccacctgtgctgc  
61 cctcgggcgccctggctggccccgcgcagagcggcgggcgctcgctgtcactgccgga  
121 ggtgagagcgcagcagtagcttcagcctgtcttgggcttgggtccagattcgctcctctgg  
181 ggctacgtcccggggaagaggaagcgaggattttgctgggggtggggctgtacctctaac  
241 agcaggtgcgcgcgaggggtgtgaacgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtgtg  
301 taagacctgcatgacgacgaggaggaacaagtgggacggcgagtgtgtcagggccag  
361 cagcaacgcatggggcgagcttcagtgtcgccagcagtgaccacaggtacggtatctact  
421 tcccagagcgcctggccgagaaataggaaagggcagccagtaggcaggccaataccca  
481 acaaaagtagaatcgagacgccctgagttcagaagttcttgaggccaaatctggctccta  
541 aaaaacatcaaaggaagcttgacacaaactctcttcagggcgcctcagaagcctgccat  
601 caccactgtgtggtgcacaatggcgccccccacaggtgtgtctctcttcattgctgtgc  
661 tgggtgacaattgcagtttgcttatgggtggatgcactcatggcaaaaaaatcactgggtgag  
1 M T R L G W P S P C C A R K  
721 catcattttaagaagacccATGACTAGACTGGGCTGGCCGAGCCCATGTTGTGCCCGTAAG  
15 Q C S E G R T Y S N A V I S P N L E T T  
781 CAGTGCAGCGAGGGGAGGACATATTCCAATGCAGTCATTTACCTAACTTGGAAACCACC  
35 R I M R V S H T F P V V D C T A A C C D  
841 AGAATCATGCGGGTGTCTCACACCTTCCCTGTCGTAGACTGCACGGCCGCTTGCTGTGAC  
55 L S S C D L A W W F E G R C Y L V S C P  
901 CTGTCCAGCTGTGACCTGGCCTGGTGGTTTCGAGGGCCGCTGCTACCTGGTGAGCTGCCCC  
75 H K E N C E P K K M G P I R S Y L T F V  
961 CACAAAGAGAACTGTGAGCCCAAGAAGATGGGCCCCATCAGGTCTTATCTCACTTTTGTG  
95 L R P V Q R P A Q L L D Y G D M M L N R  
1021 CTCCGGCCTGTTTCAGAGGCCTGCACAGCTGCTGGACTATGGGGACATGATGCTGAACAGG  
115 G S P S G I W G D S P E D I R K D L P F  
1081 GGCTCCCCCTCGGGGATCTGGGGGGACTCACCTGAGGATATCAGAAAGGACTTGCCCTTT  
135 L G K D W G L E E M S E Y S D D Y R E L  
1141 CTAGGCAAAGATTGGGGCCTAGAGGAGATGTCTGAGTACTCAGATGACTACCGGGAGCTG

155 E K D L L Q P S G K Q E P R G S A E Y T  
1201 GAGAAGGACCTCTTGCAACCCAGTGGCAAGCAGGAGCCAGAGGGAGTGCCGAGTACACG  
175 D W G L L P G S E G A F N S S V G D S P  
1261 GACTGGGGCCTACTGCCGGGCAGCGAGGGGGCCTTCAACTCCTCTGTTGGAGACAGTCCT  
195 A V P A E T Q Q D P E L H Y L N E S A S  
1321 GCGGTGCCAGCGGAGACGCAGCAGGACCCTGAGCTCCATTACCTGAATGAGTCGGCTTCA  
215 T P A P K L P E R S V L L P L P T T P S  
1381 ACCCCTGCCCCAAACTCCCTGAGAGAAGTGTGTTGCTTCCCTTGCCGACTACTCCATCT  
235 S G E V L E K E K A S Q L Q E Q S S N S  
1441 TCAGGAGAGGTGTTGGAGAAAGAAAAGGCTTCTCAGCTCCAGGAACAATCCAGCAACAGC  
255 S G K E V L M P S H S L P P A S L E L S  
1501 TCTGGAAGAGGTTCTAATGCCTTCCCATAGTCTTCCCTCCGGAAGCCTGGAGCTCAGC  
275 S V T V E K S P V L T V T P G S T E H S  
1561 TCAGTCACCGTGGAGAAAAGCCCAGTGCTCACAGTCACCCCGGGGAGTACAGAGCACAGC  
295 I P T P P T S A A P S E S T P S E L P I  
1621 ATCCCAACACCTCCCCTAGCGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATA  
315 S P T T A P R T V K E L T V S A G D N L  
1681 TCTCTACCACTGCTCCCAGGACAGTGAAAGAACTTACGGTATCGGCTGGAGATAACCTA  
335 I I T L P D N E V E L K A F V A P A P P  
1741 ATTATAACTTTACCCGACAATGAAGTTGAACTGAAGGCCTTTGTTGCGCCAGCGCCACCT  
355 V E T T Y N Y E W N L I S H P T D Y Q G  
1801 GTAGAAACAACCTACAACCTATGAATGGAATTTAATAAGCCACCCACAGACTACCAAGGT  
375 E I K Q G H K Q T L N L S Q L S V G L Y  
1861 GAAATAAAACAAGGACACAAGCAAACTCTTAACCTCTCTCAATTGTCCGTCGGACTTTAT  
395 V F K V T V S S E N A F G E G F V N V T  
1921 GTCTTCAAAGTCACCTGTTTCTAGTGAAAACGCCTTTGGAGAAGGATTGTCAATGTCACT  
415 V K P A R R V N L P P V A V V S P Q L Q  
1981 GTTAAGCCTGCCAGAAGAGTCAACCTGCCACCTGTAGCAGTTGTTTCTCCCCAACTGCAA  
435 E L T L P L T S A L I D G S Q S T D D T  
2041 GAGCTCACTTTGCCTTTGACGTCAGCCCTCATTGATGGCAGCCAAAGTACAGATGATACT  
455 E I V S Y H W E E I N G P F I E E K T S  
2101 GAAATAGTGAGTTATCATTGGGAAGAAATAAACGGGGCCCTTCATAGAAGAGAAGACTTCA  
475 V D S P V L R L S N L D P G N Y S F R L  
2161 GTTGACTCTCCCGTCTTACGCTTGCTAACCTTGATCCTGGTAACTATAGTTTCAGGTTG  
495 T V T D S D G A T N S T T A A L I V N N  
2221 ACTGTTACAGACTCGGACGGAGCCACTAACTCTACAACCTGCAGCCCTAATAGTGAACAAT  
515 A V D Y P P V A N A G P N H T I T L P Q  
2281 GCTGTGGACTACCCACCAAGTTGCTAATGCAGGACCAAATCACACCATAACTTTGCCCCAA  
535 N S I T L N G N Q S S D D H Q I V L Y E  
2341 AACTCCATCACTTTGAATGGAACAGAGCAGTGACGATCACCAGATTGTCCTCTATGAG  
555 W S L G P G S E G K H V V M Q G V Q T P  
2401 TGGTCCCTGGGTCTTGGGAGTGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCA

575 Y L H L S A M Q E G D Y T F Q L K V T D  
2461 TACCTTCATTTATCTGCAATGCAGGAAGGAGATTATACATTTACAGCTGAAGGTGACAGAT  
595 S S R Q Q S T A V V T V I V Q P E N N R  
2521 TCTTCAAGGCAACAGTCTACTGCTGTGGTGACTGTGATTGTCCAGCCTGAAAACAATAGA  
615 P P V A V A G P D K E L I F P V E S A T  
2581 CCTCCAGTGGCTGTGGCCGGCCCTGATAAAGAGCTGATCTTCCCAGTGGAAGTGCTACC  
635 L D G S S S S D D H G I V F Y H W E H V  
2641 CTGGATGGGAGCAGCAGCAGCGATGACCACGGCATTGTCTTCTACCACTGGGAGCACGTC  
655 R G P S A V E M E N I D K A I A T V T G  
2701 AGAGGCCCCAGTGCAGTGGAGATGGAAAATATTGACAAAGCAATAGCCACTGTGACTGGT  
675 L Q V G T Y H F R L T V K D Q Q G L S S  
2761 CTCCAGGTGGGGACCTACCACTTCCGTTTGACAGTGAAAGACCAGCAGGGACTGAGCAGC  
695 T S T L T V A V K K E N N S P P R A R A  
2821 ACGTCCACCCTCACTGTGGCTGTGAAGAAGGAAAATAATAGTCTCCAGAGCCCGGGCT  
715 G G R H V L V L P N N S I T L D G S R S  
2881 GGTGGCAGACATGTTCTTGTGCTTCCCAATAATTCCATTACTTTGGATGGTTCAAGGTCT  
735 T D D Q R I V S Y L W I R D G Q S P A A  
2941 ACTGATGACCAAAGAATTGTGTCCTATCTGTGGATCCGGGATGGCCAGAGTCCAGCAGCT  
755 G D V I D G S D H S V A L Q L T N L V E  
3001 GGAGATGTCATCGATGGCTCTGACCACAGTGTGGCTCTGCAGCTTACGAATCTGGTGGAG  
775 G V Y T F H L R V T D S Q G A S D T D T  
3061 GGGGTGTACACTTTCCACTTGCGAGTCACCGACAGTCAGGGGGCCTCGGACACAGACACT  
795 A T V E V Q P D P R K S G L V E L T L Q  
3121 GCCACTGTGGAAGTGCAGCCAGACCCTAGGAAGAGTGGCCTGGTGGAGCTGACCCGTCAG  
815 V G V G Q L T E Q R K D T L V R Q L A V  
3181 GTTGGTGTGGGCAGCTGACAGAGCAGCGGAAGGACACCCTTGTGAGGCAGCTGGCTGTG  
835 L L N V L D S D I K V Q K I R A H S D L  
3241 CTGCTGAACGTGCTGGACTCGGACATTAAGGTCCAGAAGATTCTGGGCCCCACTCGGATCTC  
855 S T V I V F Y V Q S R P P F K V L K A A  
3301 AGCACCGTGATTGTGTTTTATGTACAGAGCAGGCCGCCTTTCAAGGTTCTCAAAGCTGCT  
875 E V A R N L H M R L S K E K A D F L L F  
3361 GAAGTGGCCCGAAATCTGCACATGCGGCTCTCAAAGGAGAAGGCTGACTTCTTGCTTTTC  
895 K V L R V D T A G C L L K C S G H G H C  
3421 AAGGTCTTGAGGGTTGATACAGCAGGTTGCCTTCTGAAGTGTCTGGCCATGGTCACTGC  
915 D P L T K R C I C S H L W M E N L I Q R  
3481 GACCCCTCACAAAGCGCTGCATTTGCTCTCACTTATGGATGGAGAACCTTATACAGCGT  
935 Y I W D G E S N C E W S I F Y V T V L A  
3541 TATATCTGGGATGGAGAGAGCAACTGTGAGTGGAGTATATTCTATGTGACAGTGTGGCT  
955 F T L I V L T G G F T W L C I C C C K R  
3601 TTTACTCTTATTGTGCTAACAGGAGGTTTCACTTGGCTTTGCATCTGCTGCTGCAAAAGA  
975 Q K R T K I R K K T K Y T I L D N M D E  
3661 CAAAAAAGGACTAAAATCAGGAAAAAAACAAAGTACACCATCCTGGATAACATGGATGAA

995 Q E R M E L R P K Y G I K H R S T E H N  
3721 CAGGAAAGAATGGAAGTGGAGGCCAAATATGGTATCAAGCACCGAAGCACAGAGCACAAAC  
1015 S S L M V S E S E F D S D Q D T I F S R  
3781 TCCAGCCTGATGGTATCCGAGTCTGAGTTTGACAGTGACCAGGACACAATCTTCAGCCGA  
1035 E K M E R G N P K V S M N G S I R N G A  
3841 GAAAAGATGGAGAGAGGGAATCCAAAGGTTTCCATGAATGGTTCCATCAGAAATGGAGCT  
1055 S F S Y C S K D R \*  
3901 TCCTTCAGTTATTGCTCAAAGGACAGATAAtggcgcagttcattgtaaagtggaaggacc  
3961 ccttgaatccaagaccagtcagtgaggagttacagcacaaaaccactcttttagaatagt  
4021 tcattgaccttcttccccagtgaggtagatgtgtatccccacgtactaaaagaccggttt  
4081 ttgaaggcacaaaacaaaaactttgctcttttaactgagatgcttgtaataagaaataaa  
4141 ggctgggtaaaactctaaggtatataacttaaaagagttttgagtttttgtagctggcaca  
4201 atctcatattaaagatgaacaacgatttctatctgtagaaccttagagaaggtgaatgaa  
4261 acaaggttttaaaaagggtatgatttctgtcttagccgctgtgattgcctctaaggaacag  
4321 cattctaaacacggtttctctttaggacctgcagtcagatggctgtgtatgttaaaata  
4381 gcttgtctaagaggcacgggccatctgtggaggtacggagtccttgcagtagcaagcttt  
4441 ctgtgctgacggcaacactcgcacagtgccaagccctcctgggttttaattctgtgctat  
4501 gtcaatggcagttttcatctctctcaagaaagcagctgttgccattcaagagctaagga  
4561 agaatcgtattctaaggactgaggcaatagaaagggaggaggagcttaatgccgtgcag  
4621 gttgaaggtagcattgtaacattatcttttcttctctaagaaaaactacactgactcct  
4681 ctoggtgttgttttagcagtagttctctaagttaaaccggatccccagtttacattaat  
4741 gcaatagaagtgattaattcattaagcatttattatgttctgtaggctgtgcgtttggac  
4801 tgccatagatagggataacgactcagcaattgtgtatatattccaaaactctgaaataca  
4861 gtcagtccttaacttggtatggcgtggttatgatactctgggtccccgacagggtactttccaa  
4921 aataacttgacatagatgtattcaactcatatgtttaaaaatacatttaagtttttctac  
4981 cgaataaatcttattttcaaacatgaaagacaattaaaacattcccaccacacaaagcagta  
5041 ctcccagcaattaaactggaggttaattgtagcctgctacgttgactgggtcagggtagtt  
5101 ccccatccacccttggtcctgaggctgggtggccttggtggtgcccttggcattttttgtg  
5161 ggaagattagaatgagagatagaaccagtggtgtgtacctaaagtgtgagcacacctaaac  
5221 aatatcctgttgacaaatgcttttttaacacatgggaaaactaggaatgcattgctgatg  
5281 aagaagcaaggtattttaaacaccagggcaggagtgccagagaaaatgtttccccatgggt  
5341 tcttaaaaaaattcagcttttaggtgcttttgcacatctccggagtagttcatcctcatg  
5401 ggaccatcttatttttacttattgtaatttactggggaaaggcagaactaaaaagtgtgt  
5461 cattttatttttaaaataattgctttgcttatgcctacactttctgtataactagccaat  
5521 tcaatactgtctatagtgttagaaggaatgtgatttttttttaaccagttattgag  
5581 cttcataagcctagaatctgccttatcaggtgaccagggttatgggtgtttgcatgcaaa  
5641 tgtgaatttctggcataggggacagcagcccaatgtaaagtcacgggcgtaatgagga  
5701 agaagggagtgaaacatttaccgctttatgtacataacatatgcagtttacatactcattt  
5761 gatccttataatcaaccttgaagaggagatactatcattcttatgttgcatagccctc  
5821 tgaagggccagagaggttaagtaacttcccagaggtcatggccaagaagtagtggctcca  
5881 agaactgaatgcaaatttttttaactgtagagttctgctttccactaaacaaagaactcc  
5941 tgccttgatggatggaggggcaaattctggtggaacttttgggccacctgaaagttctatt

```

6001 cccaggactaagaggaatttcttttaattggatccagagagccaaggtcagagggagagat
6061 ggcttgcatagtctcctgtggatcacacccgggccacccctccctctaggtttacagtgg
6121 acttcttctgcccctcctccttttctgtccttggccatctcagcctggcctctctgatcc
6181 ttccatcacagaaggatcttgaatctctgggaaatcaaacatcacagtagtgatcagaaa
6241 gtgagtcctgtcttgtcaccccatcttctcatcagaacaaagcacgagatggaatgaccaa
6301 ccagcattcttcatggtggactgcttatcattgaggatcttgggagataaagcacgcta
6361 agagctctggacagagaaaaacaggccctagaatatgggagtgggtgtttgtagggtctca
6421 taggctaacaagcacttttagttgctggtttacattcaatgaaggaggattcatacccatg
6481 gcattacaaggctaagcatgtgtatgactaaggaactatctgaaaaacatgcagcaaggt
6541 aagaaaatgtaccactcaacaagccagtgtatgccaccttttgtgcgcggggaggagagtg
6601 actaccattgttttttgtgtgacaaagctatcatggactattttaattcttggttttattg
6661 cttaaaatatattatttttccctatgtgttgacaagggtatttctaatatcacactattaa
6721 atatatgcactaatctaaataaagggtgtctgtattttctgtaatgcttatttttaggggg
6781 aaatttgttttctttatgcttcagggtagaggattcccttgagtataggtcagcaaaact
6841 ctggcctgcagcctgtgtgtgcacgccccatgagccgaaaagtgggtcttatgttttcaa
6901 atgggttaaaaataaataaaaaaatttgaacatgtgaactatatgacattcagattttgtg
6961 ttcataaataaagttttattggaacatatcc

```

**Figure 2D. 254P1D6B v.4 through v.20, SNP variants of 254P1D6B v.1.** The 254P1D6B v.4 through v.20 proteins have 1072 amino acids. Variants 254P1D6B v.4 through v.20 are variants with single nucleotide difference from 254P1D6B v.1. 254P1D6B v.5 and v.6 proteins differ from 254P1D6B v.1 by one amino acid. 254P1D6B v.4 and v.7 through v.20 proteins code for the same protein as v.1. Though these SNP variants are shown separately, they can also occur in any combinations and in any of the transcript variants listed above in Figures 2A, Figure 2B and Figure 2C.

Variant	Nucleic acid position	Nucleic Acid Variation	Amino Acid Position	Amino Acid Variation
254P1D6B v.4	286	C/G	Silent variant	
254P1D6B v.5	935	C/A	142	P=>T
254P1D6B v.6 (Identical AA as v.2)	980	T/G	157	S=>A
254P1D6B v.7	2347	G/A	Silent variant	
254P1D6B v.8	3762	C/T	Silent variant	
254P1D6B v.9	3772	A/G	Silent variant	
254P1D6B v.10	3955	C/T	Silent variant	
254P1D6B v.11	4096	C/T	Silent variant	
254P1D6B v.12	4415	G/A	Silent variant	
254P1D6B v.13	4519	G/A	Silent variant	
254P1D6B v.14	4539	A/G	Silent variant	

Variant	Nucleic acid position	Nucleic Acid Variation	Amino Acid Position	Amino Acid Variation
254P1D6B v.15	4614	G/T	Silent variant	
254P1D6B v.16	5184	G/C	Silent variant	
254P1D6B v.17	5528	T/G	Silent variant	
254P1D6B v.18	5641	G/A	Silent variant	
254P1D6B v.19	6221	T/C	Silent variant	
254P1D6B v.20	6223	G/A	Silent variant	

**Figure 3:**

**Figure 3A. Amino acid sequence 254P1D6B v.1 clone LCP-3 (SEQ ID NO.: 8).** The 254P1D6B v.1 clone LCP-3 protein has 1072 amino acids.

```
1 MAPPTGVLSS LLLLVTIAGC ARKQCSEGR T YSNAVISP NL ETTRIMRVSH TFPVVDCTAA
61 CCDLSSCDLA WWFEGRCYL V SCPHKENCEP KKMGP IRSYL TFVLRPVQRP AQLLDYGDM
121 LNRGSPSGIW GDSPEDIRK D LPFLGKDWGL EEMSEYSDDY RELEKDLLQP SGKQEPGSA
181 EYTDWGLLP G SEGAFNSSVG DSPAVPAETQ QDPELHYLNE SASTPAPKLP ERSVLLPLPT
241 TPSSGEVLEK EKASQLQE QS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST
301 EHSIPTPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIEE
481 KTSVDSPVLR LSNLDPGNYS FRLTVTDS DG ATNSTTAALI VNNAV DYPPV ANAGPNHTIT
541 LPQNSITLNG NQSSDDHQIV LYEWSLGPGS EGKHVVMQGV QTPYLHLSAM QEGDYTFQLK
601 VTDSSRQQST AVVTVIVQPE NNRPPVAVAG PDKELIFPVE SATLDGSSSS DDHGIVFYHW
661 EHVRGPSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR
721 ARAGGRHVLV LPNNSITLDG SRSTDDQ RIV SYLWIR DGQS PAAGDVIDGS DHSVALQLTN
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF
901 LLFKVLRVDT AGCLLKCSGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT
961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST
1021 EHNSSLMVSE SEFSDSDQTI FSREKMERGN PKVSMNGSIR NGASFSYCSK DR
```

**Figure 3B. Amino acid sequence 254P1D6B v.2 (SEQ ID NO.: 9).** The 254P1D6B v.2 protein has 1072 amino acids.

```
1 MAPPTGVLSS LLLLVTIAGC ARKQCSEGR T YSNAVISP NL ETTRIMRVSH TFPVVDCTAA
61 CCDLSSCDLA WWFEGRCYL V SCPHKENCEP KKMGP IRSYL TFVLRPVQRP AQLLDYGDM
121 LNRGSPSGIW GDSPEDIRK D LPFLGKDWGL EEMSEYADDY RELEKDLLQP SGKQEPGSA
181 EYTDWGLLP G SEGAFNSSVG DSPAVPAETQ QDPELHYLNE SASTPAPKLP ERSVLLPLPT
241 TPSSGEVLEK EKASQLQE QS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST
301 EHSIPTPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIEE
481 KTSVDSPVLR LSNLDPGNYS FRLTVTDS DG ATNSTTAALI VNNAV DYPPV ANAGPNHTIT
541 LPQNSITLNG NQSSDDHQIV LYEWSLGPGS EGKHVVMQGV QTPYLHLSAM QEGDYTFQLK
601 VTDSSRQQST AVVTVIVQPE NNRPPVAVAG PDKELIFPVE SATLDGSSSS DDHGIVFYHW
661 EHVRGPSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR
721 ARAGGRHVLV LPNNSITLDG SRSTDDQ RIV SYLWIR DGQS PAAGDVIDGS DHSVALQLTN
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF
901 LLFKVLRVDT AGCLLKCSGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT
```



961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSQDTI FSREKMERGN PKVSMNGSIR NGASFSYCSK DR

**Figure 3C. Amino acid sequence 254P1D6B v.3 (SEQ ID NO: 10).** The 254P1D6B v.3 protein has 1063 amino acids.

1 MTRLGWSPSPC CARKQCSEGR TYSNAVISPN LETTRIMRVS HTFPVVDCTA ACCDLSSCDL  
61 AWWFEGRCYL VSCPHKENCE PKKMGPIRSY LTFVLRPVQR PAQLLDYGDM MLNRGSPSGI  
121 WGDSPEDIRK DLPFLGKDWG LEEMSEYSDD YRELEKDLLQ PSGKQEPGRS AEYTDWGLLP  
181 GSEGAFNSSV GDSPAVPAET QQDPELHYLN ESASTPAPKL PERSVLLPLP TTPSSGEVLE  
241 KEKASQLQEQ SSNSSGKEVL MPShSLPPAS LELSSVTVEK SPVLTVTGPS TEHSIPTPPT  
301 SAAPSESTPS ELPISPTTAP RTVKELTVSA GDNLIITLPD NEVELKAFVA PAPPVETTYN  
361 YEWNLISHPT DYQGEIKQGH KQTLNLSQLS VGLYVFKVTV SSENAFGEGF VNVTVKPARR  
421 VNLPPVAVVS PQLQELTLPL TSALIDGSQS TDDTEIVSYH WEEINGPFIE EKTSVDSFVL  
481 RLSNLDPGNY SFRLTVTDSG GATNSTTAAL IVNNAVDPYP VANAGPNHTI TLPQNSITLN  
541 GNQSSDDHQI VLYEWSLPGG SEGKHVVMQG VQTPYLHLSA MQEGDYTFQL KVTDSRQOS  
601 TAVVTVIVQP ENNRPPVAVA GPKELIFPV ESATLDGSSS SDDHGIVFYH WEHVRGPSAV  
661 EMENIDKAIA TVTGLQVGTY HFRLTVKDQQ GLSSTSTLTV AVKKENNSPP RARAGGRHVL  
721 VLPNNSITLD GSRSTDDQRI VSYLWIRDGQ SPAAGDVIDG SDHSVALQLT NLVEGVYTFH  
781 LRVTD SQGAS DTDATVEVQ PDPRKSGLVE LTLQVGVGQL TEQRKDTLVR QLAVLLNVLD  
841 SDIKVQKIRA HSDLSTVIVF YVQSRPPFKV LKAAEVARNL HMRLSKEKAD FLLFKVLRVD  
901 TAGCLLKCSG GHGCDPLTKR CICSHLWMEN LIQRYIWDGE SNCEWSIFYV TVLAFTLIVL  
961 TGGFTWLCIC CCKRQKRTKI RKKTKYTILD NMDEQERMEL RPKYGIKHRS TEHNSSLMVS  
1021 ESEFDSQDT IFSREKMERG NPKVSMNGSI RNASFSYCS KDR

**Figure 3D. Amino acid sequence 254P1D6B v.5 (SEQ ID NO: 11).** The 254P1D6B v.5 protein has 1072 amino acids.

1 MAPPTGVLSS LLLLVTIAGC ARKQCSEGRT YSNAVISPNL ETTRIMRVSH TFPVVDCTAA  
61 CCDLSSCDLA WWFEGRCYLV SCPHKENCEP KKMGPISYSL TFVLRPVQRP AQLLDYGDM  
121 LNRGSPSGIW GDSPEDIRK LTFGLGKDWGL EEMSEYSDDY RELEKDLLQP SGKQEPGRSA  
181 EYTDWGLLPG SEGAFNSSVG DSPAVPAETQ QDPELHYLNE SASTPAPKL PERSVLLPLPT  
241 TPSSGEVLEK EKASQLQEQS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST  
301 EHSIPTPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP  
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV  
421 NVTVPKARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIEE  
481 KTSVDSFVLR LSNLDPGNYS FRLTVTDSG ATNSTTAALI VNNAVDPYPV ANAGPNHTIT  
541 LPQNSITLNG NQSSDDHQIV LYEWSLPGGS EGKHVVMQGV QTPYLHLSAM QEGDYTFQLK  
601 VTDSSRQOST AVVTVIVQPE NNRPPVAVAG PDKELIFPVE SATLDGSSSS DDHGIVFYHW  
661 EHVRGPSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTV VKKENNSPPR  
721 ARAGGRHVLV LPNNSITLDG SRSTDDQRIV SYLWIRDGQS PAAGDVIDGS DHSVALQLTN  
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ  
841 LAVLLNVLDs DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF  
901 LLFKVLRVDT AGCLLKCSGH GHGCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT

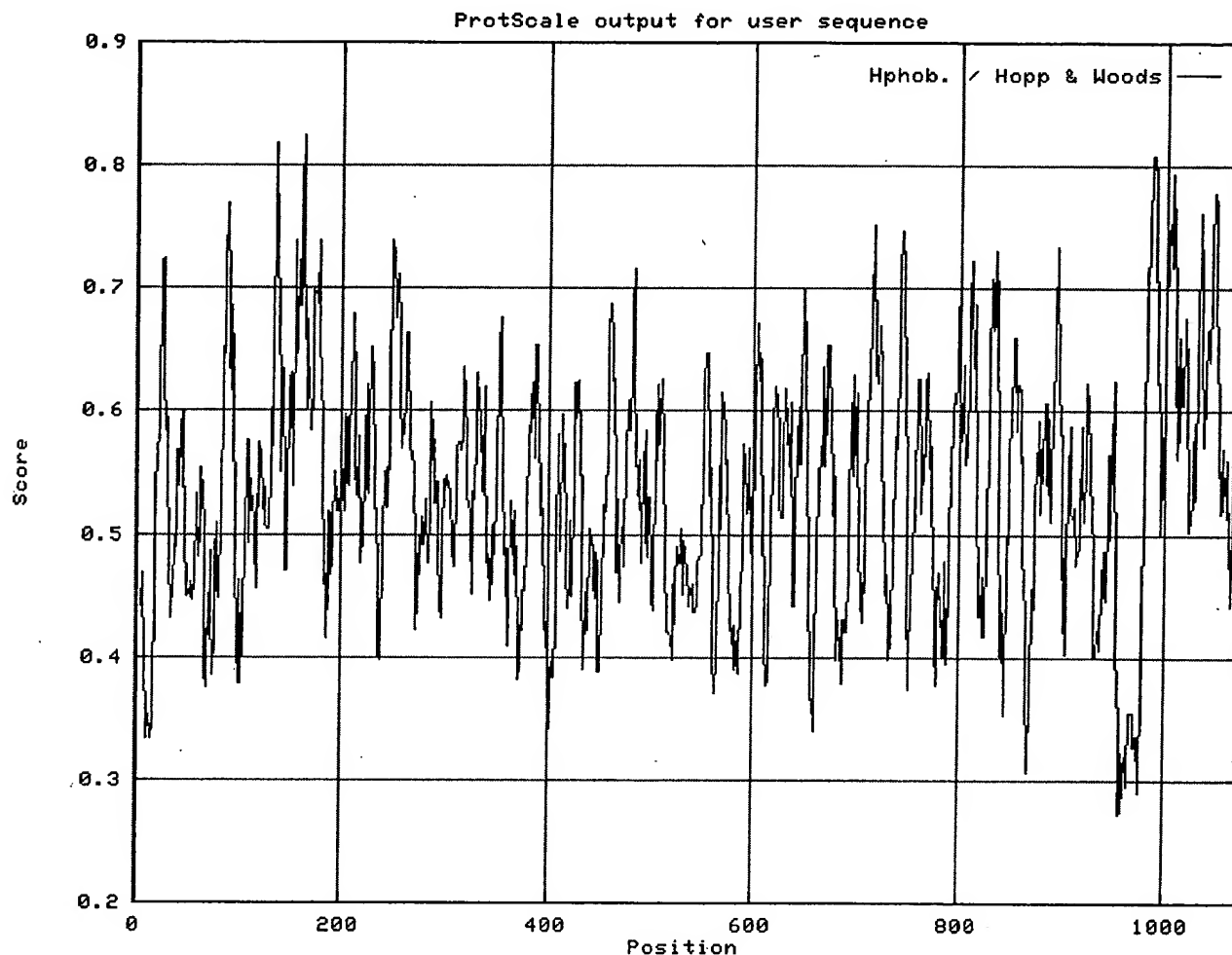
961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSQDTI FSREKMERGN PKVSMNGSIR NGASFSYCSK DR

**Figure 3E. Amino acid sequence 254P1D6B v.6 (SEQ ID NO: 12).** The 254P1D6B v.6 protein has 1072 amino acids.

1 MAPPTGVLSS LLLLVTIAGC ARKQCSEGRT YSNAVISPNI ETTRIMRVSH TFPVVDCTAA  
61 CCDLSSCDLA WWFEGRCYLV SCPHKENCEP KKMGPISYLV TFVLRPVQRP AQLLDYGDMM  
121 LNRGSPSGIW GDSPEDIRKD LPFLGKDWGL EEMSEYADDY RELEKDLLQP SGKQEPGSA  
181 EYTDWGLLPG SEGAFNSSVG DSPAVPAETQ QDPHELHYLNE SASTPAPKLP ERSVLLPLPT  
241 TPSSGEVLEK EKASQLQEQS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST  
301 EHSIPTPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP  
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV  
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIEE  
481 KTSVDSPVLR LSNLDPGNYS FRLTVTDS DG ATNSTTAALI VNNAVDPYPV ANAGPNHTIT  
541 LPQNSITLNG NQSSDDHQIV LYEWSLGPGS EGKHVVMQGV QTPYLHLSAM QEGDYTFQLK  
601 VTDSSRQQST AVVTVIVQPE NNRPPVAVAG PDKELIFPVE SATLDGSSSS DDHGIVFYHW  
661 EHVRGPSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR  
721 ARAGGRHVLV LPNNSITLDG SRSTDDQRIV SYLWIRDGQS PAAGDVIDGS DHSVALQLTN  
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ  
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF  
901 LLFKVLRVDT AGCLLKCSGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT  
961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSQDTI FSREKMERGN PKVSMNGSIR NGASFSYCSK DR



**Figure 5: 254P1D6B variant 1**  
**Hydrophilicity profile**  
(Hopp T.P., Woods K.R., 1981.  
Proc. Natl. Acad. Sci. U.S.A. 78:3824-3828)



**Figure 6: 254P1D6B variant 1**  
**Hydropathicity Profile**  
(Kyte J., Doolittle R.F., 1982. J. Mol. Biol. 157:105-132)

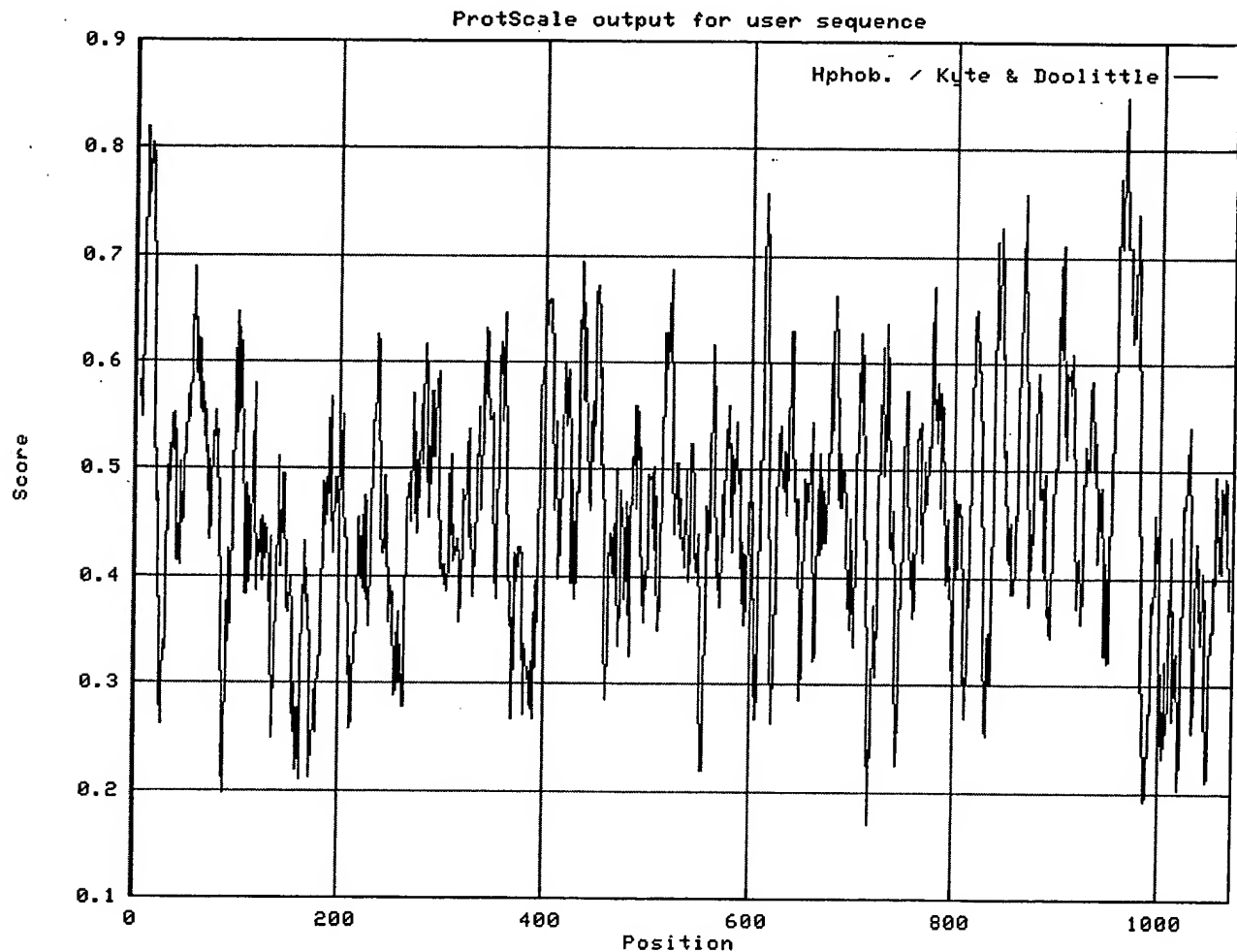
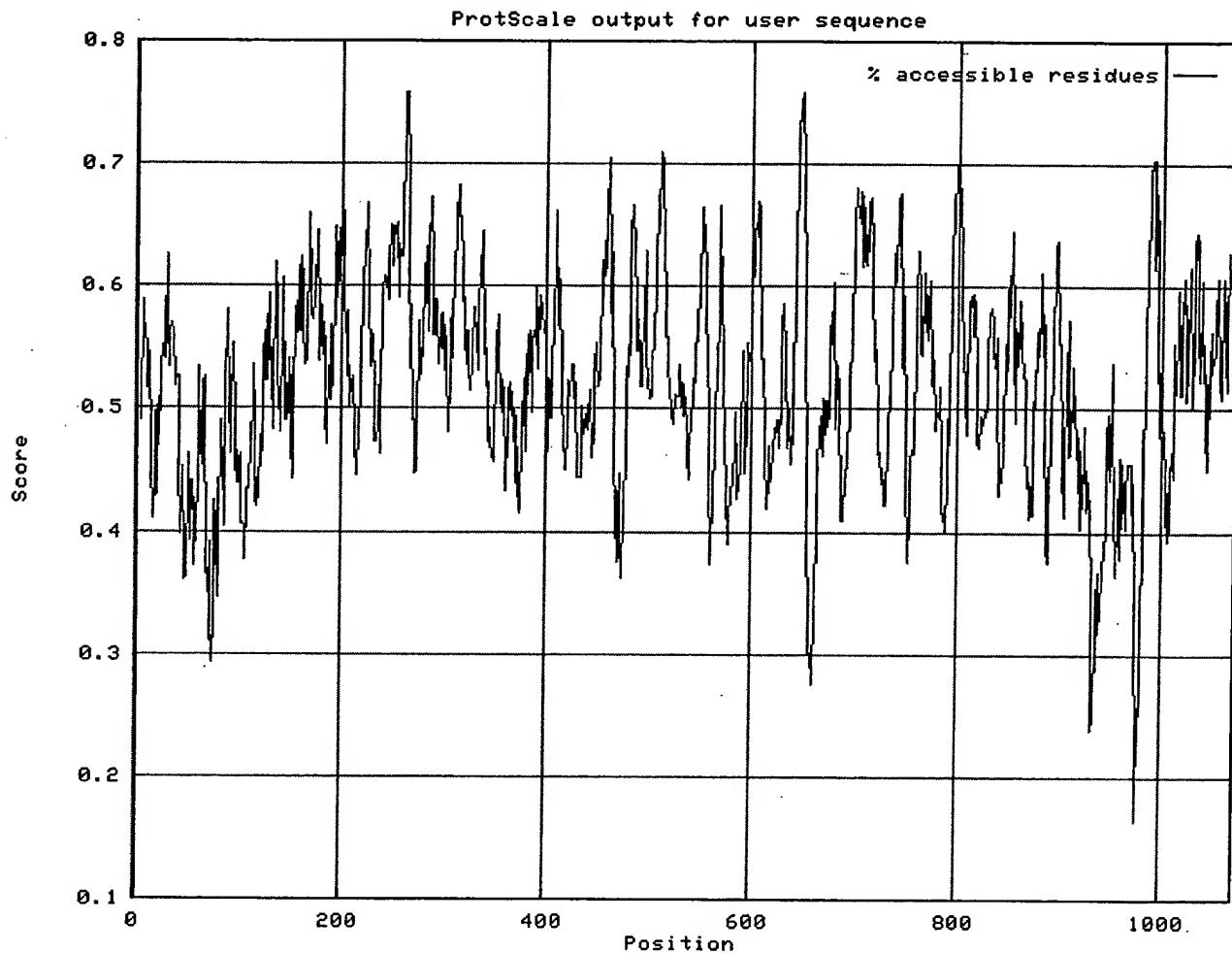
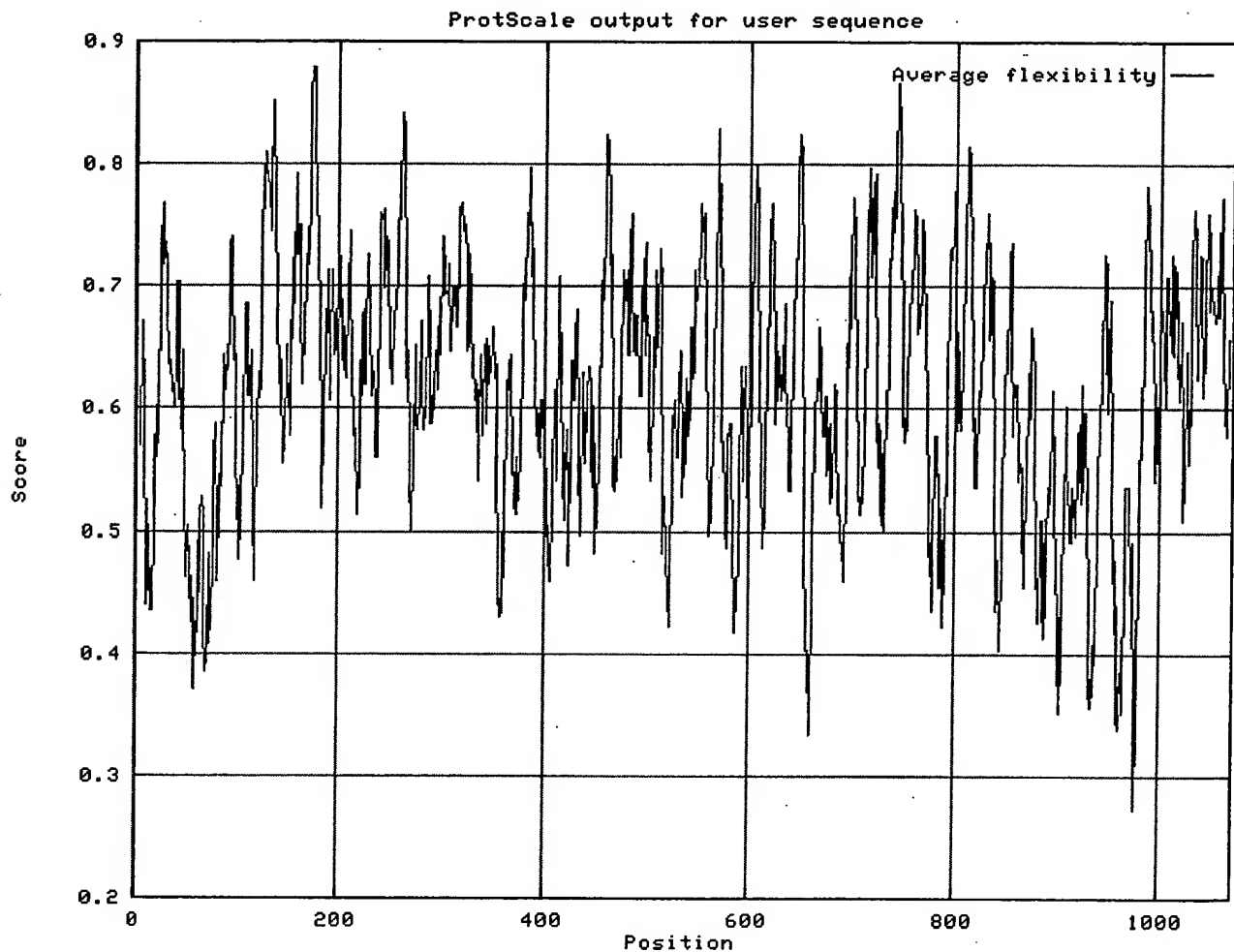


Figure 7: 254P1D6B variant 1  
% Accessible Residues Profile  
(Janin J., 1979. Nature 277:491-492)



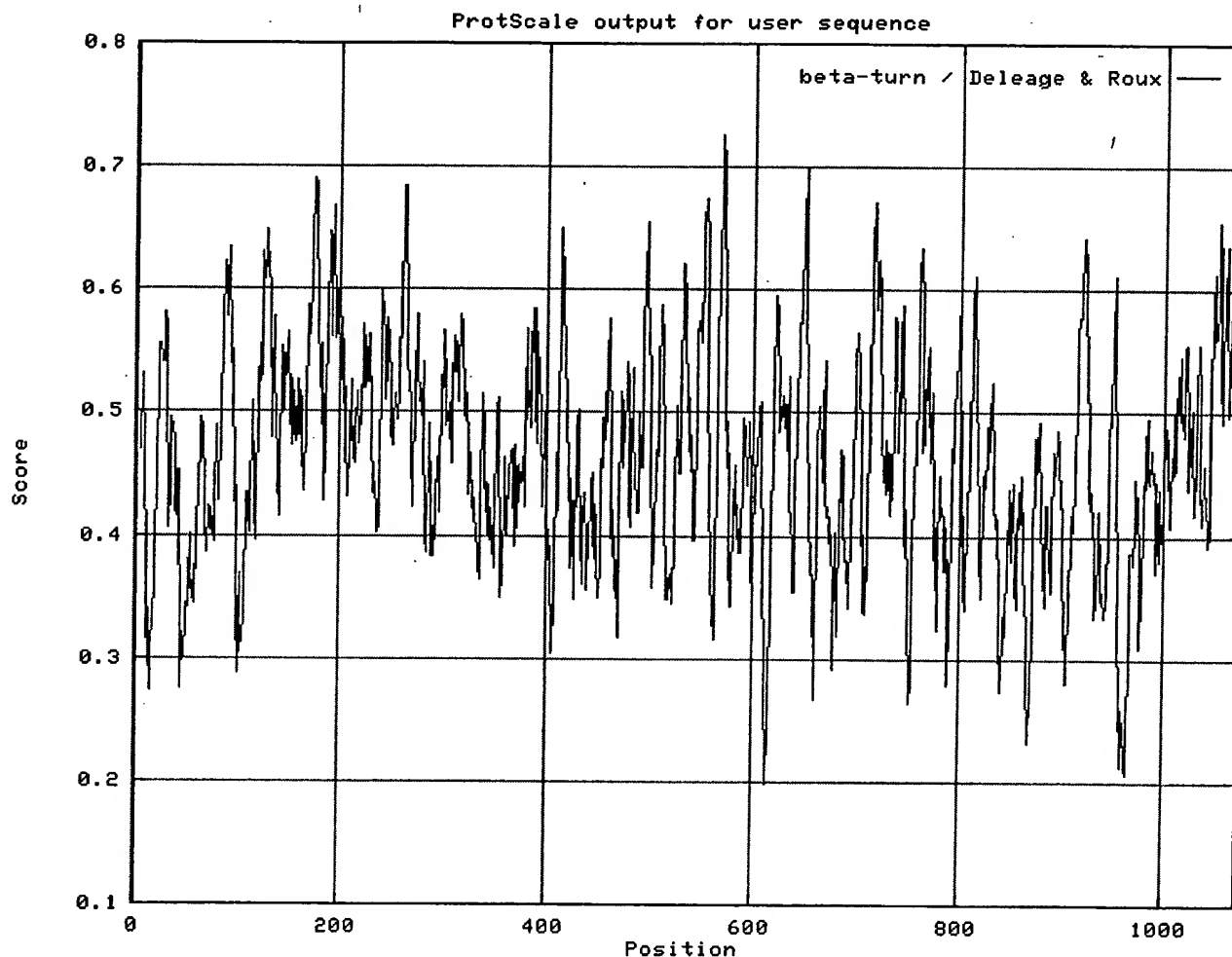
**Figure 8: 254P1D6B variant 1**  
**Average Flexibility Profile**  
(Bhaskaran R., Ponnuswamy P.K., 1988.  
Int. J. Pept. Protein Res. 32:242-255)



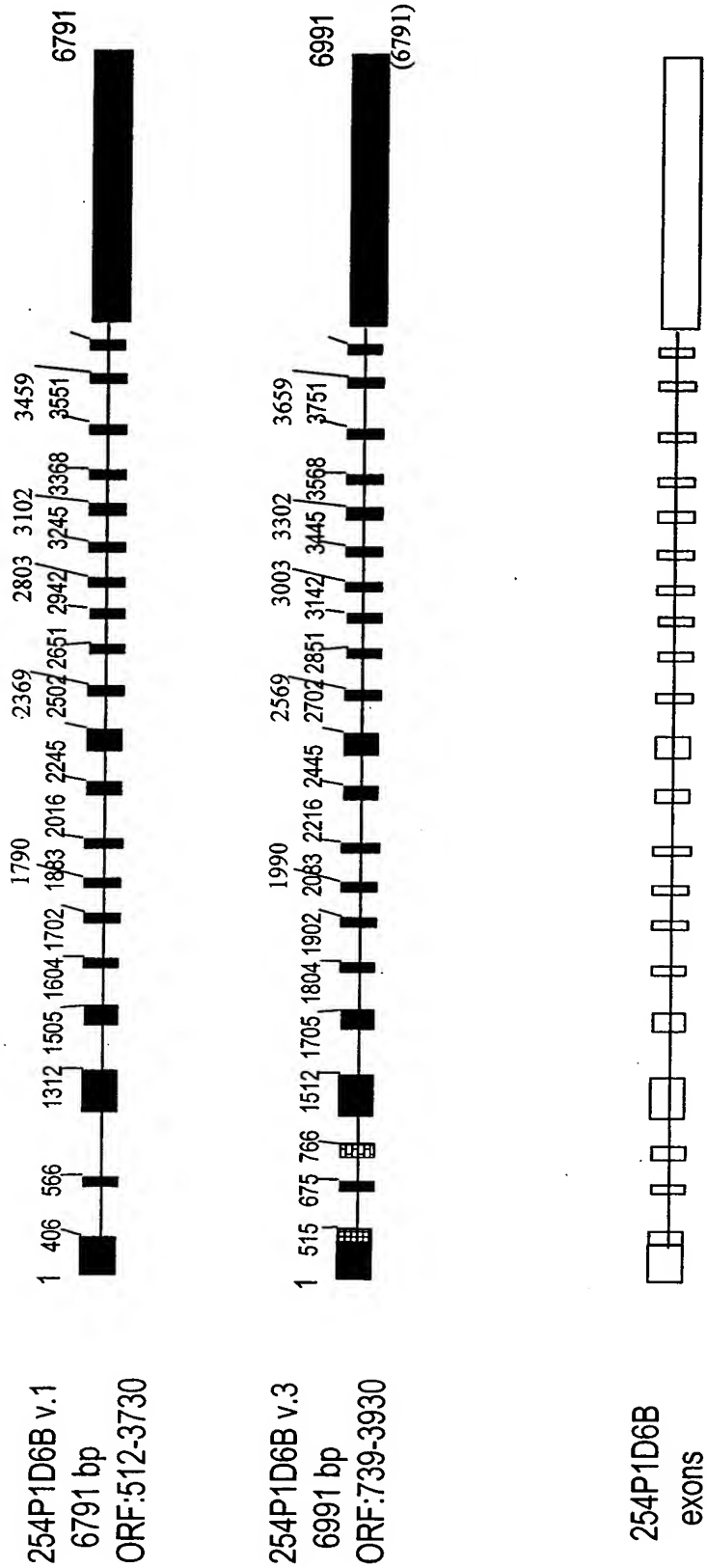
## Figure 9: 254P1D6B variant 1

### Beta-turn Profile

(Deleage, G., Roux B. 1987.  
Protein Engineering 1:289-294)

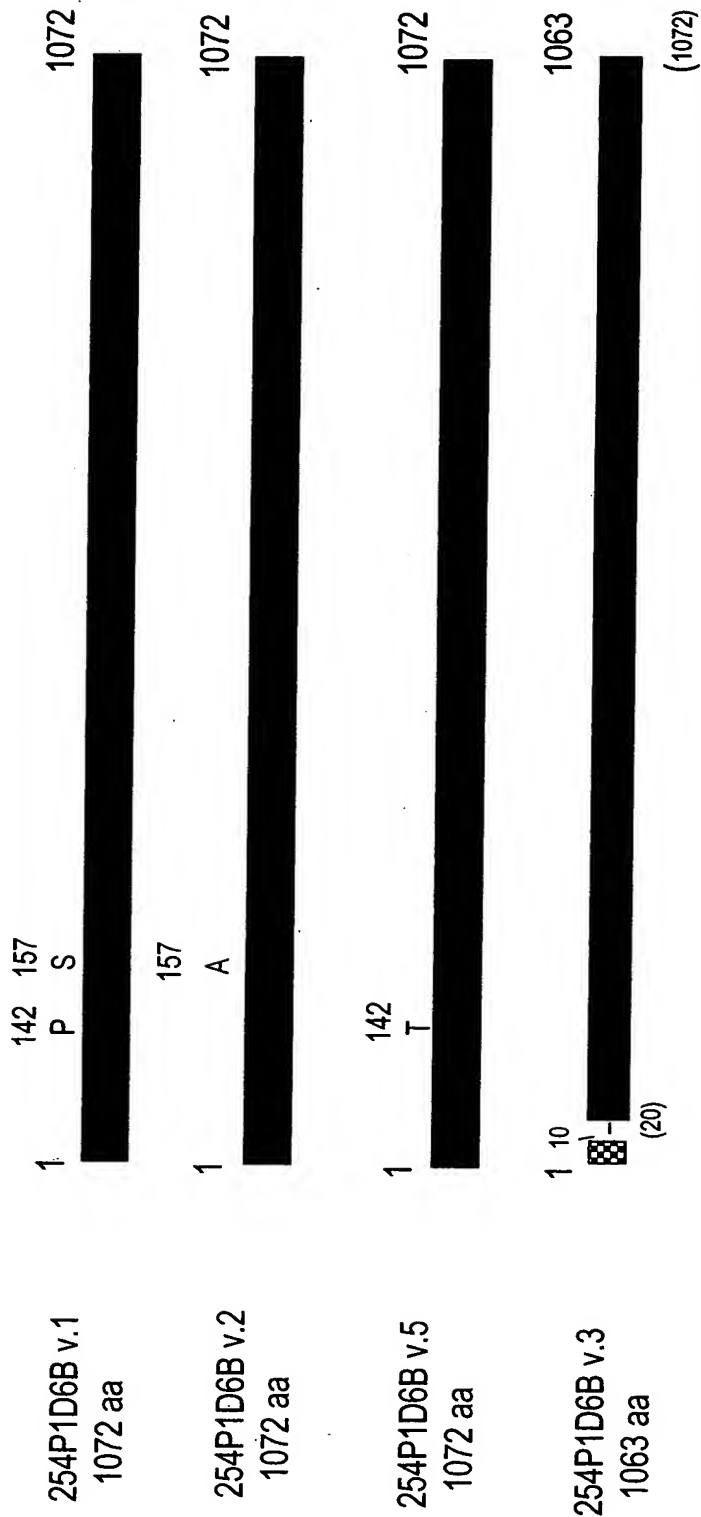


**Figure 10**





**Figure 11**



## Figure 12

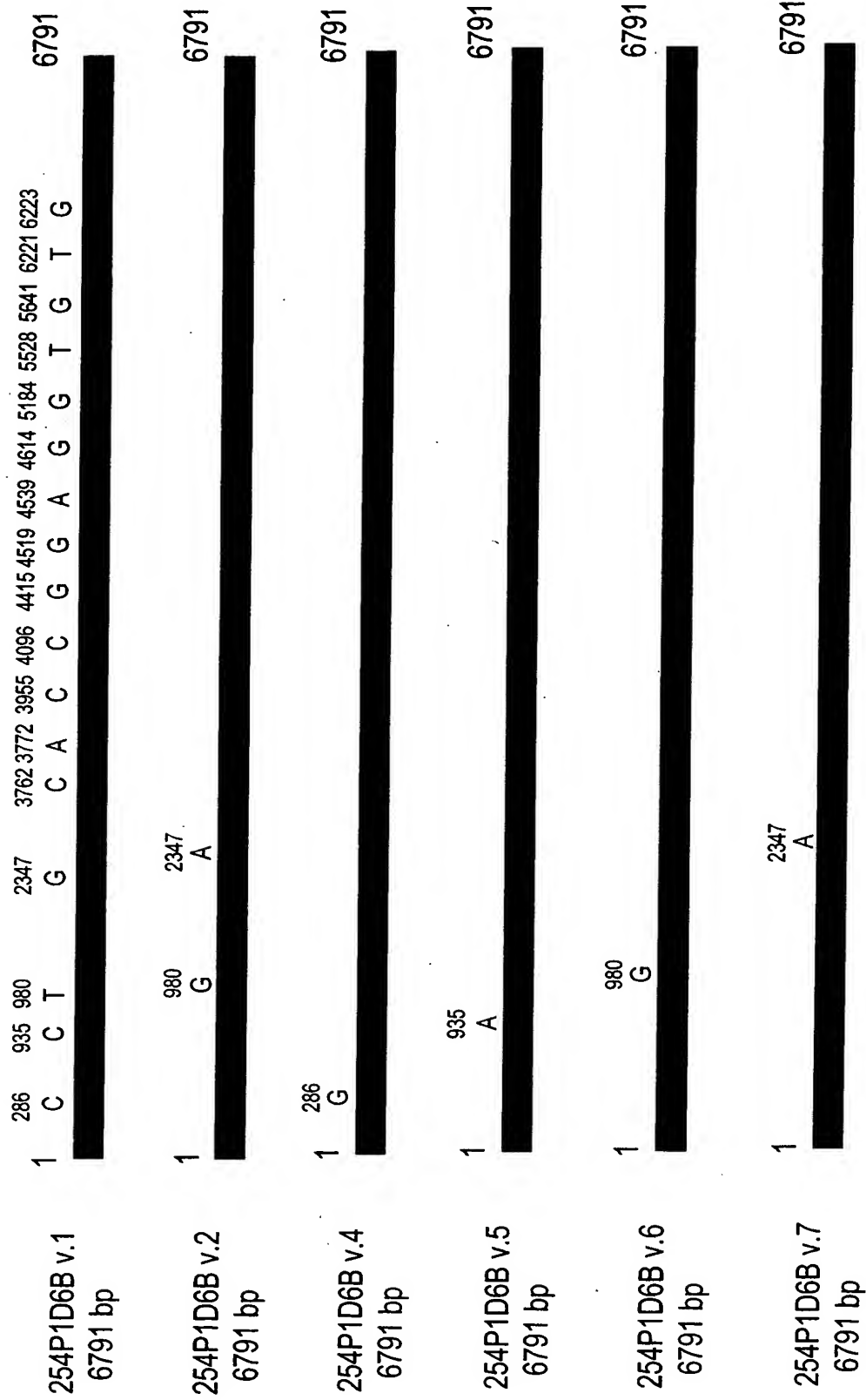


Figure 12 (con'd)

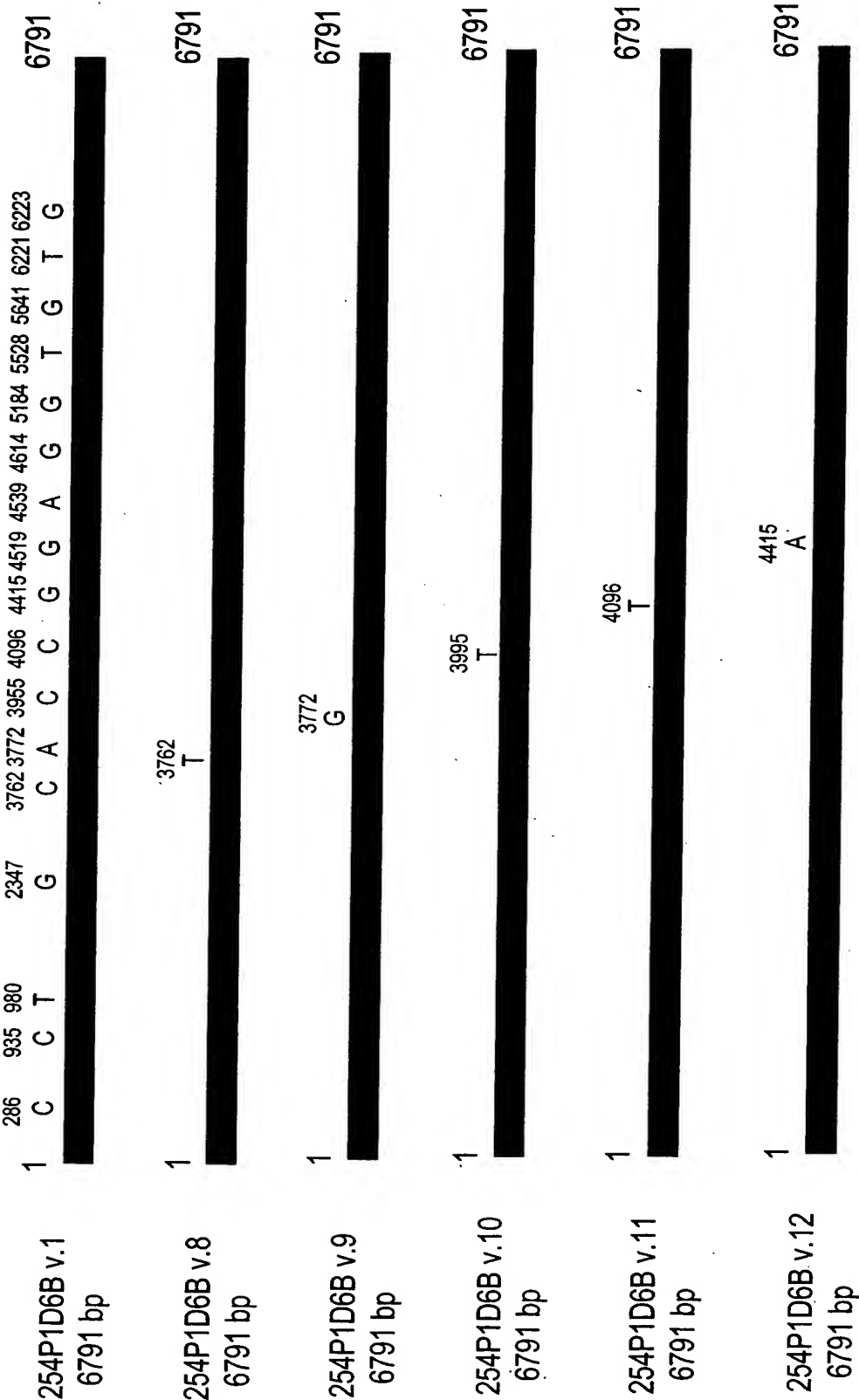


Figure 12 (con'd)

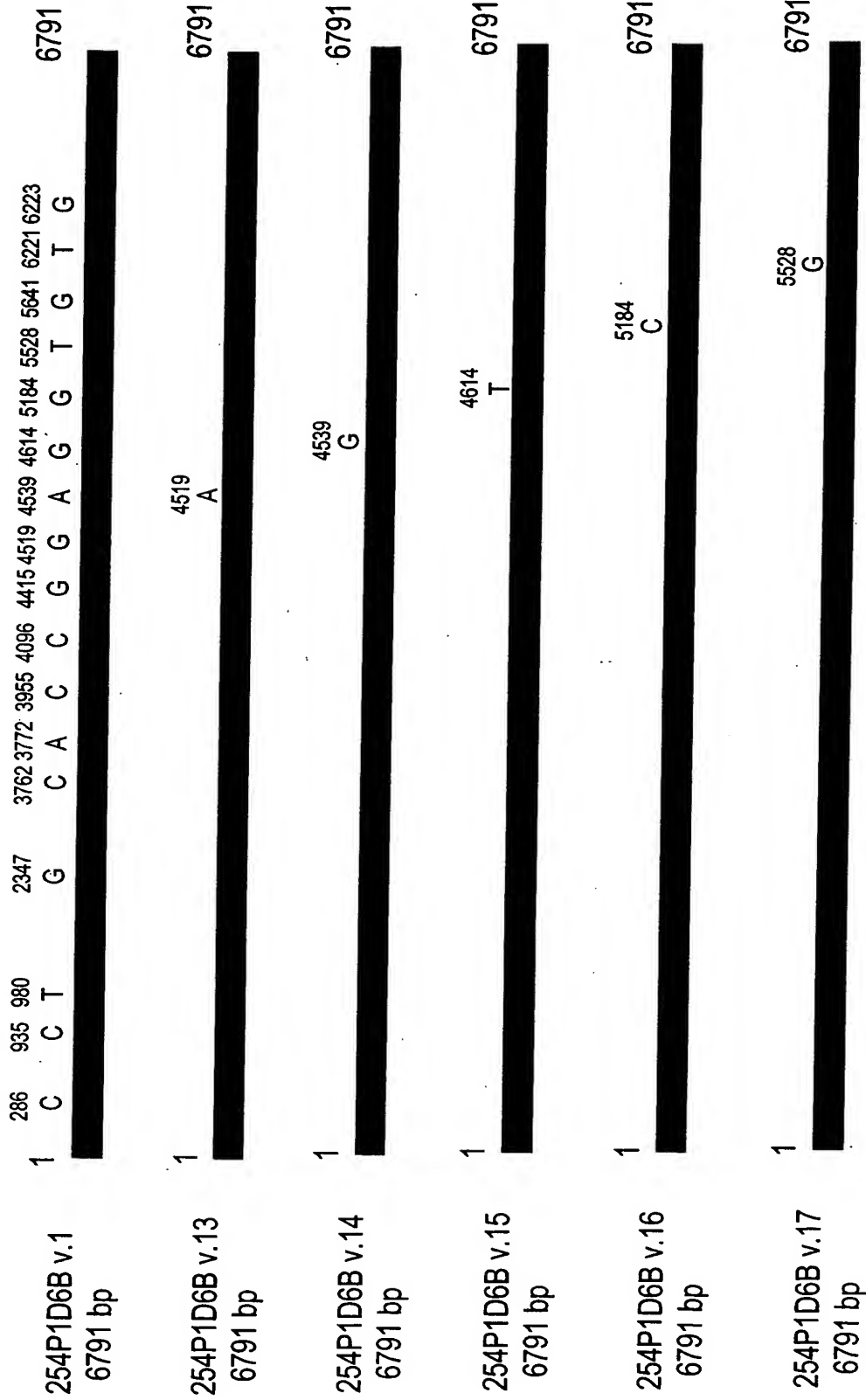
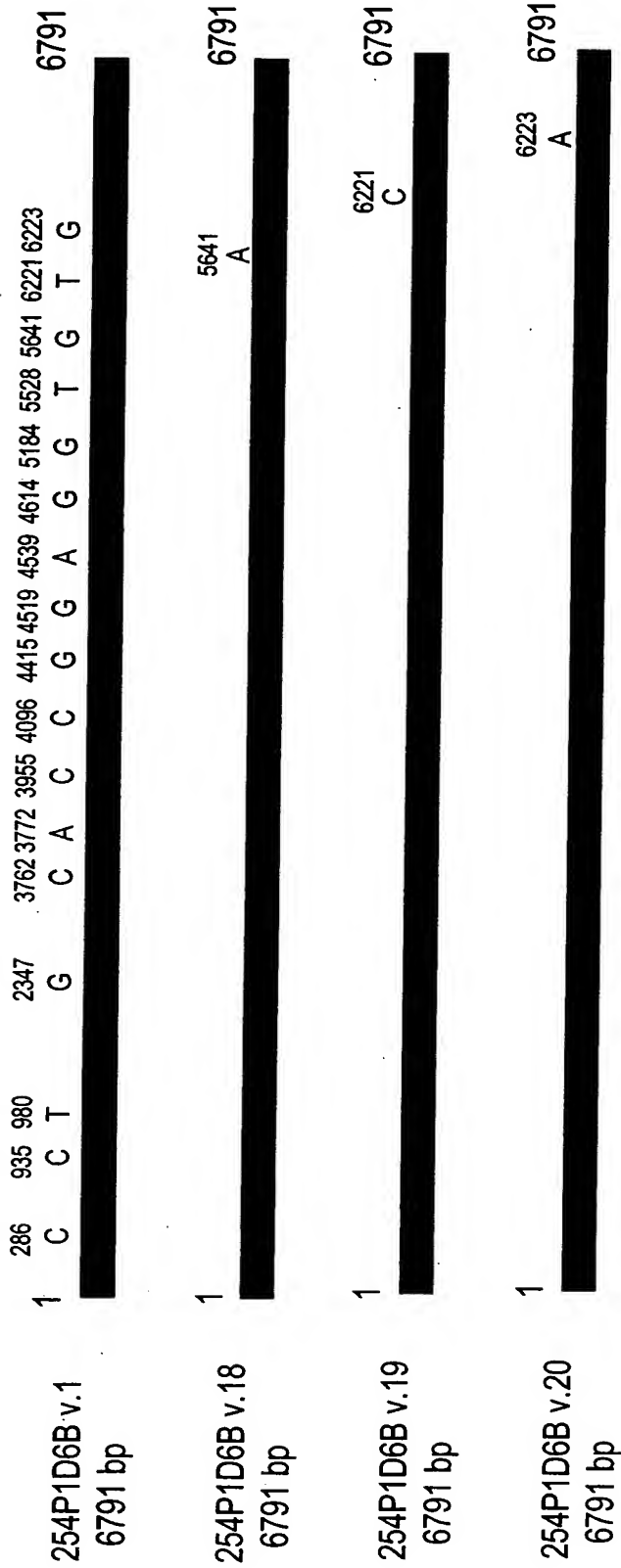
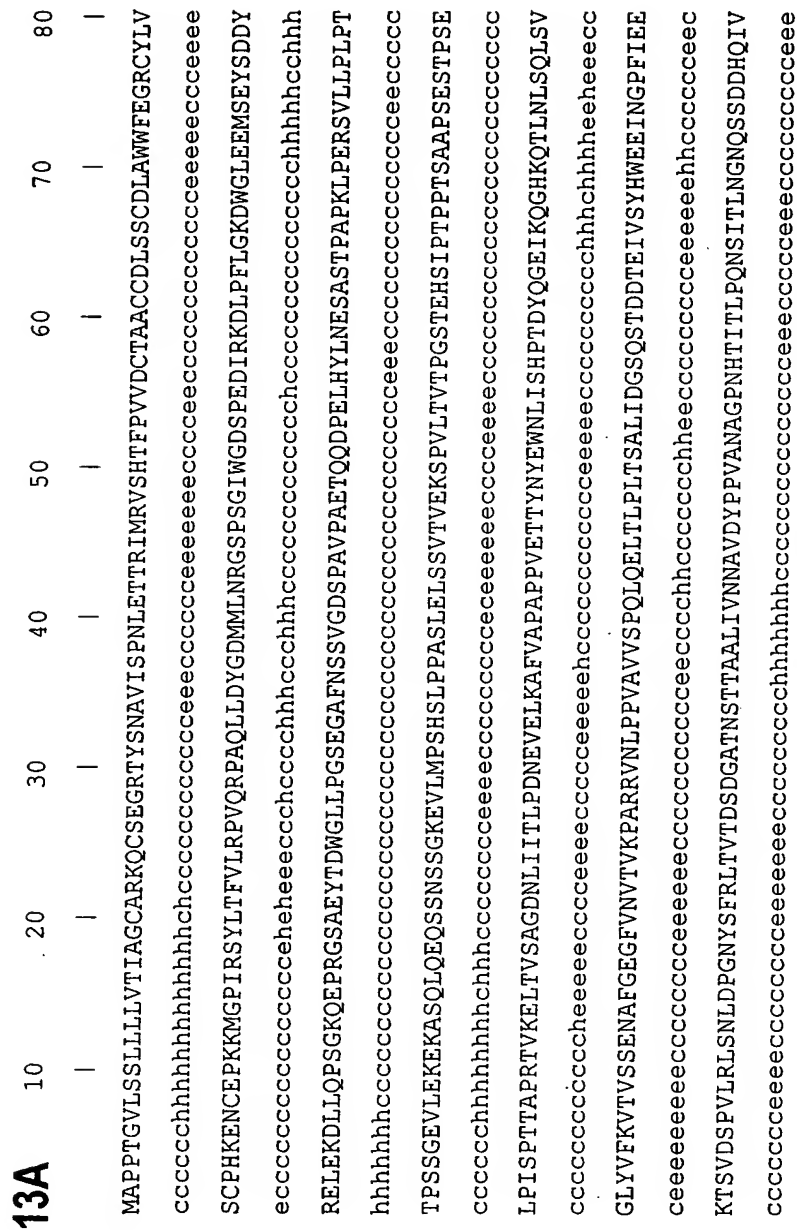


Figure 12 (con'd)



**Figure 13: Secondary structure prediction of 254P1D6B variant 1**



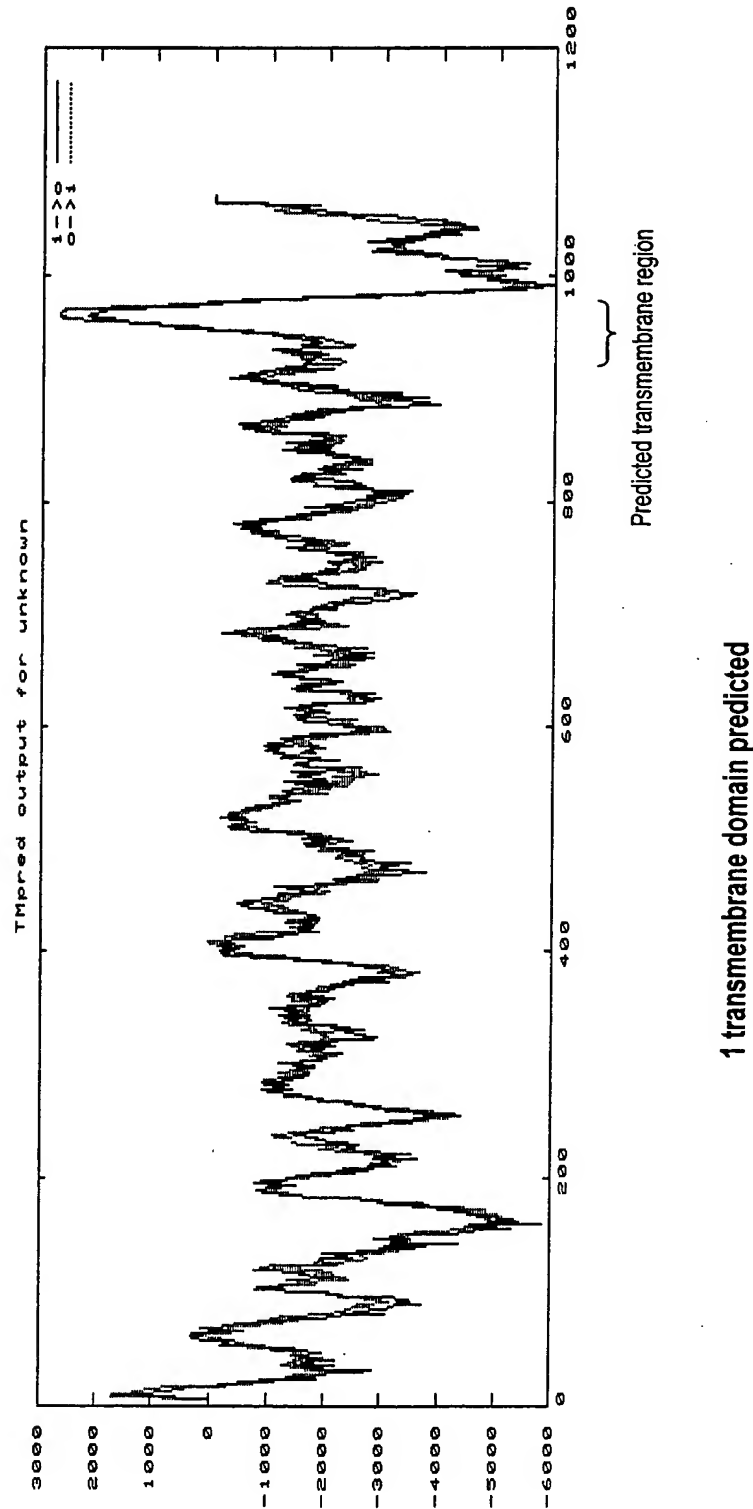
Alpha helix(h): 18.19%  
Extended strand (e): 24.81%  
Random coil(c): 57.00%

# Secondary structure prediction of 254P1D6B variant 1 (continued)

570	580	590	600	610	620	630	640						
LYEWSLPGSEGGKHVVMQGVQTPYLHLSAMQEGDYTFQLKVTDSRRQQTAVVTIVQPNRNPPVAVAGDPDKELIFPVE	eeeecc	SATLDGSSSSDDHGIVFYHWEHVRGSAVEMENIDKAIATVTGLQVGTYHFRLTVKDQQGLSSTSTLTVAVKKENNSPPR	cc	ARAGGRHVLVLPNNSITLDGSRSTDDQRIVSYLWIRDGQSPAAGDIVDGS DHSVALQLTNLVEGVYTFHLRVTDTSQGASD	cc	TDTATVEVQDPDKSLVELTLQVGVQLTEQRKDTLVRQLAVLLNVLSDIKVQKIRAHSDLSTVIVFYVQSRPPFKVL	cc	KAAEVARNLHRLSKEADFLLFKVLRVDTAGCLLKCSGHGCHCDPLTKRCICSHLWMENLIQRYIWDGESNCEWSIFYVT	hh	VLAFTLIVLTGGFTWLCICCCCKRQRTKIRKTKYTILDNMDEQERMELRPKYGIKHRSTEHNSSLVMVSESEFSDQDTI	hheeeeecc	FSREKMERGNPKVSMNGSIRNGASFYSCKDR	ehhhhhcc

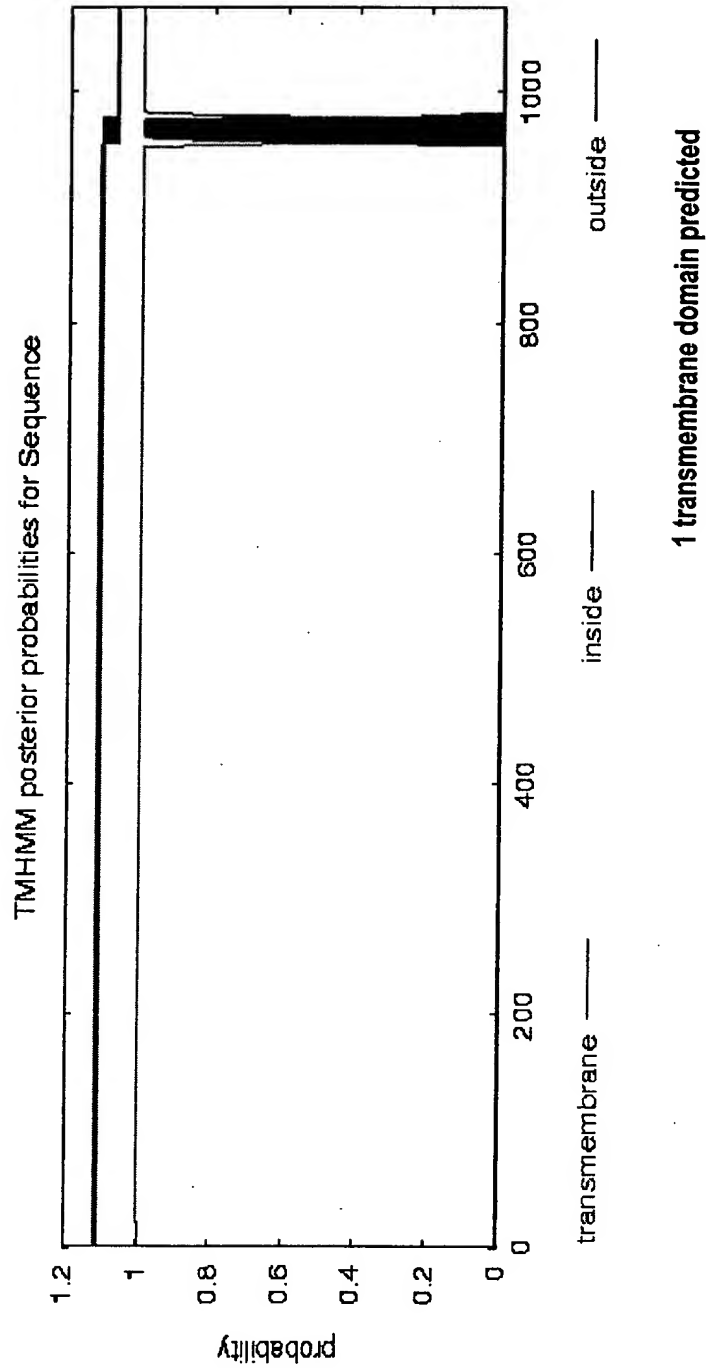
Alpha helix(h): 18.19%  
Extended strand (e): 24.81%  
Random coil(c): 57.00%

**Figure: Transmembrane prediction for 254P1D6B variant 1  
13B**



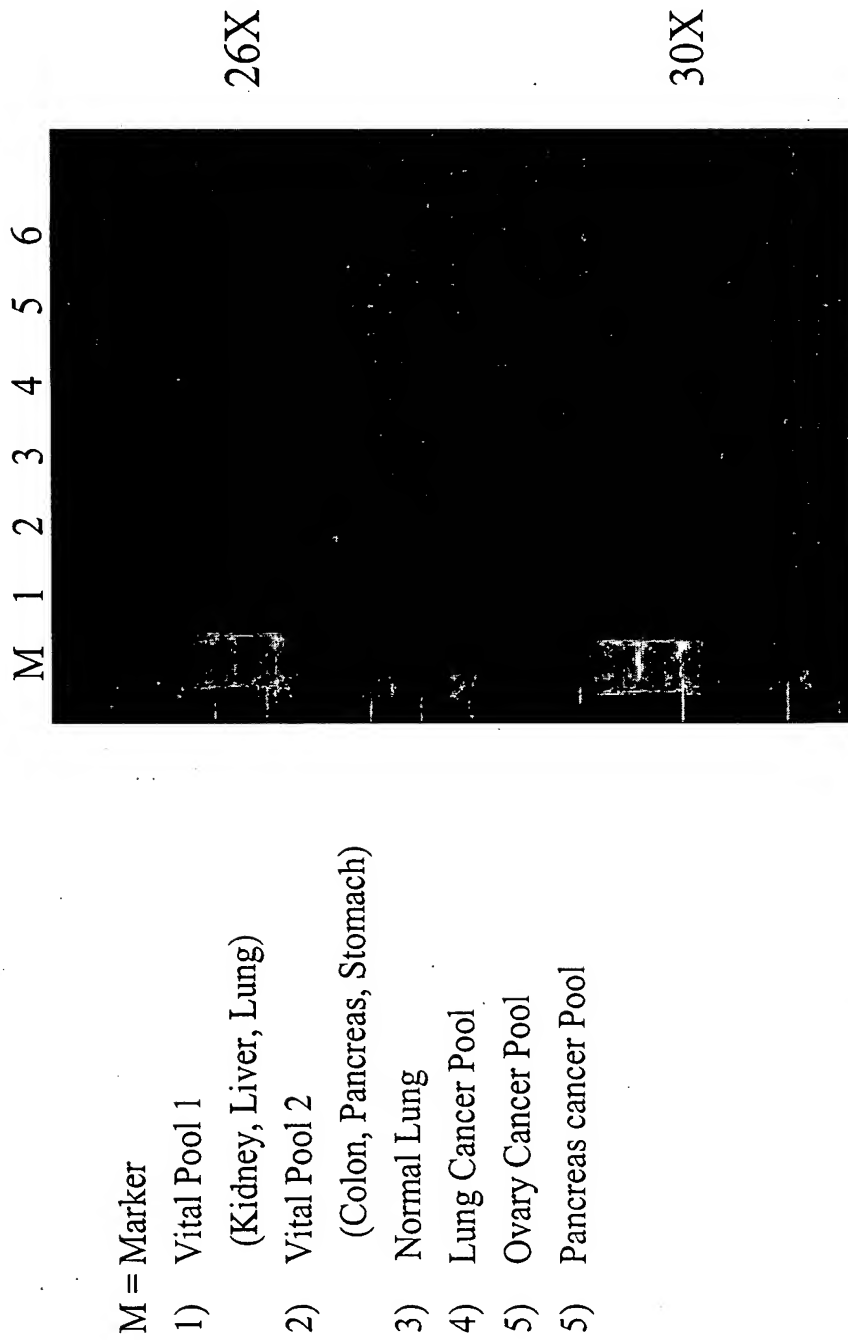


**Figure: Transmembrane prediction for 254P1D6B variant 1**  
**13C**

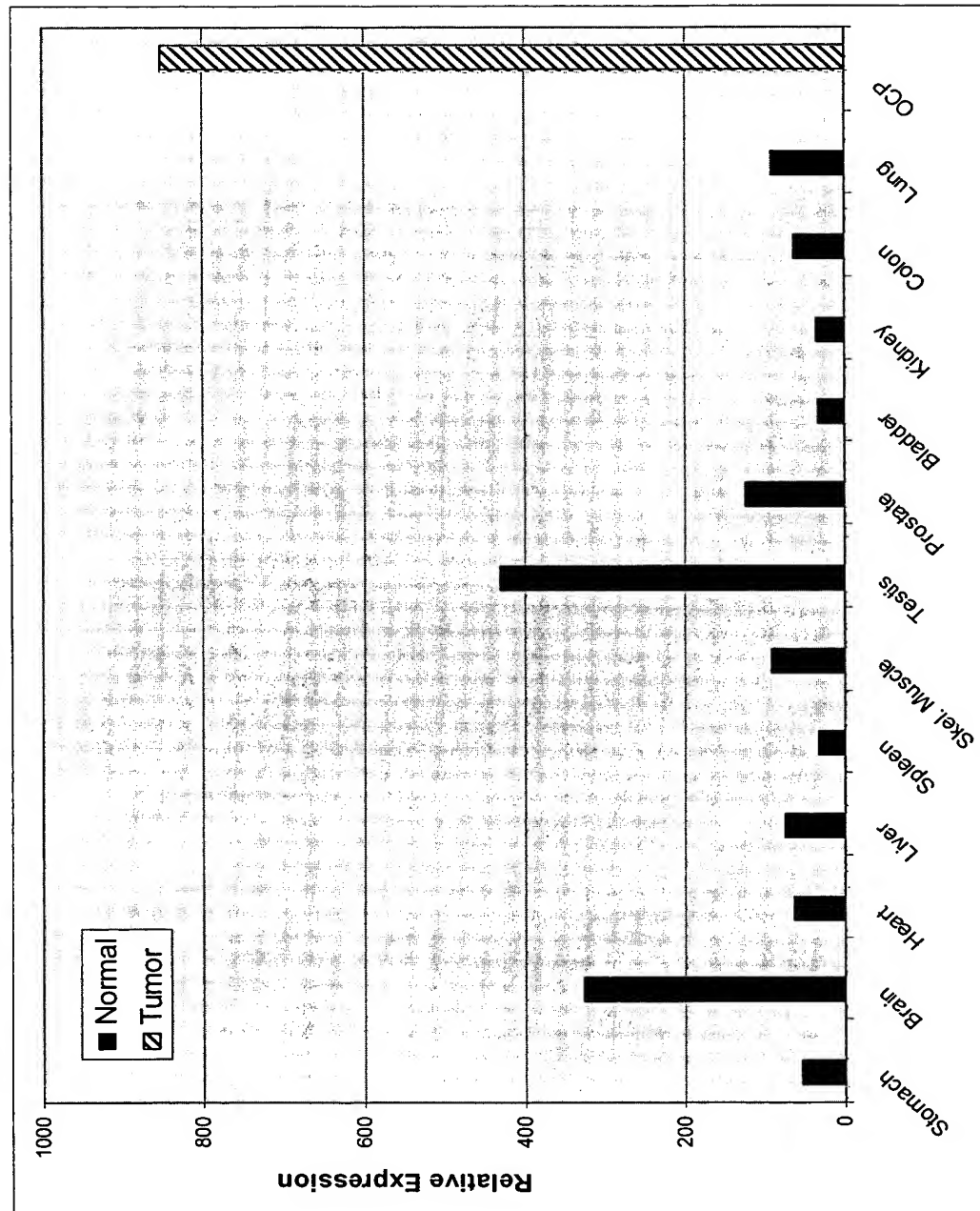


BEST AVAILABLE COPY

Figure 14A 254P1D6B Expression by RT-PCR



**Figure 14B Expression of 254P1D6B in  
Normal Human Tissues and Ovarian Cancer Patient Specimens**

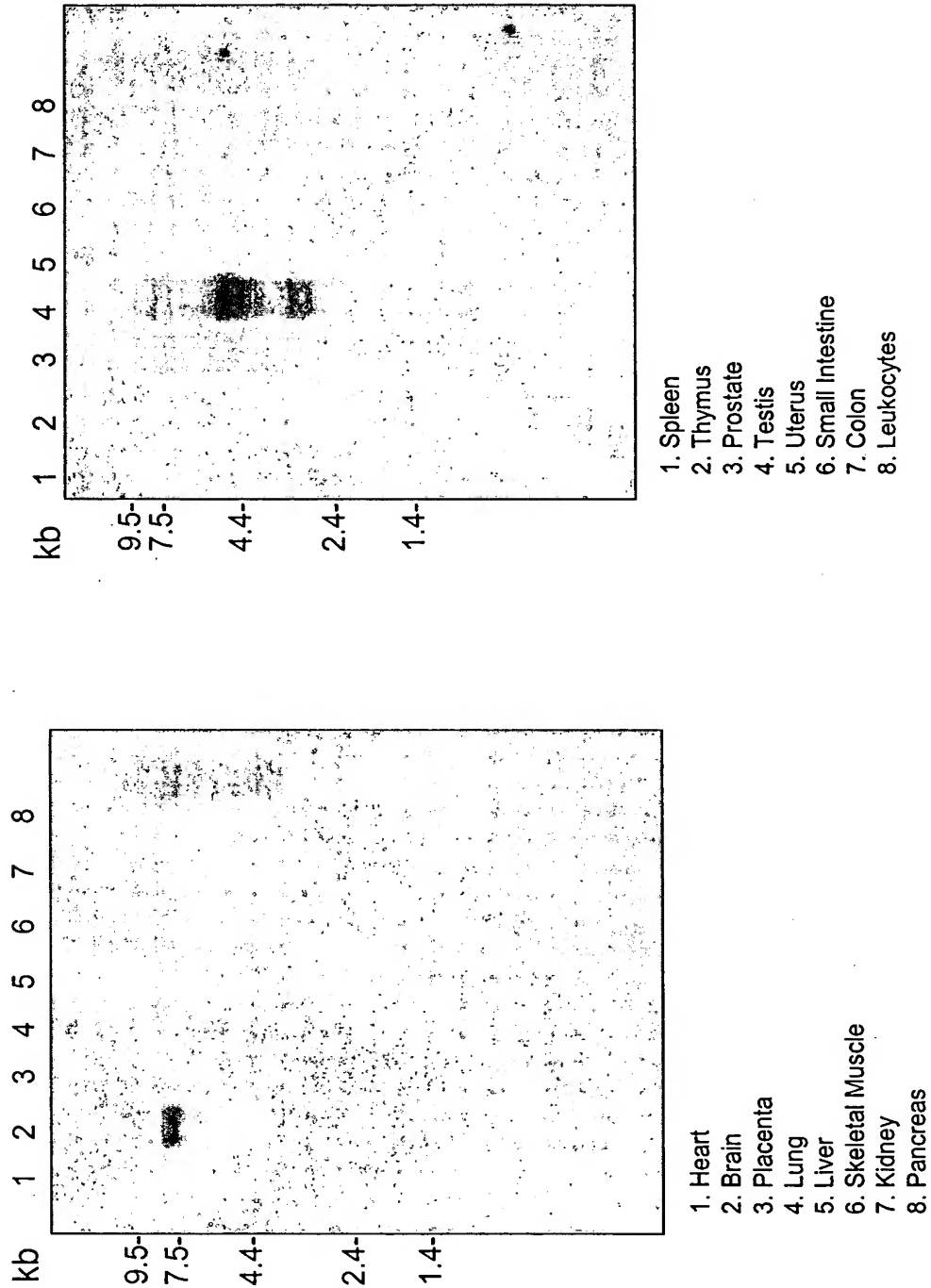


THIS PAGE BLANK (USPTO)

Sheet 36 of 38

BEST AVAILABLE COPY

**Figure 15 Expression of 254P1D6B in Normal Tissues**



**Figure 16** Expression of 254P1D6B in Lung Cancer Patient Specimens

Panel#	Pathology	Grade	Expression
1	Normal		
2	A427 Cell line		
3	Adeno	3	
4	Adeno	I	
5	Adeno	IB	
6	Adeno	IB	
7	Adeno	IIIA	
8	Adeno	IIIA	
9	Adeno	Mod Diff	
10	Adeno	Mod Diff	
11	Adeno		
12	Bronchioalv.	IA	
13	Large Cell	I	
14	Large Cell	IIB	
15	Large Cell	IIIA	
16	Large Cell	IV	
17	Papillary	I	
18	Papillary	IB	
19	Papillary	IV	
20	Small Cell	I	
21	Small Cell	I	
22	Small Cell	I	
23	Small Cell	IIB	
24	Squamous	IB	
25	Squamous	IB	
26	Squamous	IB	
27	Squamous	IIB	
28	Squamous	IIB	
29	Squamous	IIIA	
30	Squamous	IIIA	
31	Squamous		
32	Squamous		
33	Squamous		

	No Expression
	Low Expression
	High expression

Figure 17: Expression of 254P1D6b in 293T cells

